

09/508 704

Att 17

WEST Search History

DATE: Monday, March 10, 2003

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<i>DB=USPT,PGPB,JPAB,EPAB,DWPI; PLUR=YES; OP=ADJ</i>			
L8	11 with 16 with L7	972	L8
L7	virus or viral or virally or viruses	126869	L7
L6	coat protein	5429	L6
L5	comovir\$	93	L5
L4	comovirus	92	L4
L3	cpmv or (cow pea mosaic virus)	40	L3
L2	comovir?	0	L2
L1	plant	531187	L1

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Select?	Document ID	Section(s)	Page(s)	# Pages to print	Database
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<input checked="" type="checkbox"/>	5173410	all	all	19	USPT,PGPB,JPAB,EPAB,DWPI
<input checked="" type="checkbox"/>	5304730	all	all	33	USPT,PGPB,JPAB,EPAB,DWPI
<input checked="" type="checkbox"/>	JP363014693A	all	all	N/A	USPT,PGPB,JPAB,EPAB,DWPI
<input checked="" type="checkbox"/>	JP402049591A	all	all	N/A	USPT,PGPB,JPAB,EPAB,DWPI
<input checked="" type="checkbox"/>	WO8904868A	all	all	N/A	USPT,PGPB,JPAB,EPAB,DWPI
<input checked="" type="checkbox"/>	JP401120290A	all	all	N/A	USPT,PGPB,JPAB,EPAB,DWPI

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cm1



11e14



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[Generate Collection](#)[Print](#)**Search Results - Record(s) 951 through 972 of 972 returned.**

- └ 951. [ZA 9104563 A](#). Protecting plants against cucumber mosaic virus infection - relates to e.g. isolated coat protein of virus strain and isolated DNA. COLLAD, D D A, et al. A01H000/00 A01N000/00 C12N000/00.
- └ 952. [AU 9171951 A](#) [US 5824856 A](#) [CA 2037677 A](#) [JP 04121200 A](#) [AU 636717 B](#). Prod'n. of exogenous gene or its prod. in plant cells - comprises using cDNA of RNA replicase and recombinant virus genomic RNA contg. exogenous gene. FURUSAWA, I, et al. A01H001/00 A01H004/00 A01H005/00 C12N005/00 C12N005/14 C12N015/00 C12N015/10 C12N015/33 C12N015/40 C12N015/52 C12N015/82 C12P021/00 C12P021/00 C12R001:91.
- └ 953. [WO 9117254 A](#) [IL 98031 A](#) [AU 9178655 A](#) [US 5143905 A](#) [CN 1059760 A](#) [US 5306628 A](#). Extending host range or toxicity of insecticidal proteins - using protein capable of binding to gut epithelium of insects. FEDERICI, B A, et al. A01N063/00 A61K037/00 A61K039/07 A61K039/12 C07K003/08 C07K013/00 C07K019/00 C12N001/21 C12N015/00 C12N015/09 C12N015/31 C12N015/62 C12N015/63 C12P021/00 C12P021/02.
- └ 954. [WO 9108296 A](#) [DE 69014100 E](#) [EP 460217 A](#) [EP 460217 A4](#) [EP 460217 B1](#) [JP 04144685 A](#). DNA obtd. by cleavage of cDNA corresp. to RNA - of Odontoglossum ring spot virus coding for viral coat protein and is useful as probe and vector for plant gene recombination. CHATANI, M, et al. C07K013/00 C07K015/04 C12N001/21 C12N005/10 C12N015/40 C12N015/82 C12P021/02 C12Q001/68 C12R001/19 C12P021/02 C12R001:19.
- └ 955. [CA 2026703 C](#) [EP 425004 A](#) [NL 8902452 A](#) [NL 9001711 A](#) [CA 2026703 A](#) [PT 95494 A](#) [JP 03280883 A](#) [EP 425004 B1](#) [DE 69028479 E](#) [ES 2094744 T3](#) [IE 76133 B](#) [US 6093554 A](#) [US 6197542 B1](#) [CA 2396794 A1](#). Recombinant DNA comprising RNA virus derived sequences - used for protecting organisms, esp. plants from virus infection or for producing protein or RNA. AMELOOT, P, et al. A01H001/04 A01H005/00 C07G017/00 C12N001/00 C12N001/21 C12N005/00 C12N005/02 C12N005/04 C12N005/10 C12N015/00 C12N015/09 C12N015/11 C12N015/40 C12N015/79 C12N015/82 C12P019/34 C12P021/00 C12P021/04 C12P021/06.
- └ 956. [WO 9104332 A](#) [US 5677157 A](#) [AU 9062840 A](#) [EP 491733 A1](#) [JP 05500308 W](#) [EP 491733 B1](#) [DE 69014644 E](#) [ES 2065545 T3](#). Regeneration and transformation of squash plants - by transfer and integration of genetic materials into genome of squash plants. CHEE, P P. A01H001/04 A01H004/00 A01H005/00 C12N005/04 C12N015/82.
- └ 957. [US 4970168 A](#). Virus-resistant plants - obtd. by transforming plants with DNA which encodes coat proteins of Potato Viruses-X and Y. TUMER, N E. C12N001/00 C12N005/00 C12N015/00.
- └ 958. [JP 02109992 A](#). Prod'n. of useful proteins - using genes of proteins specific to infection. C12N005/14 C12N015/63 C12P021/00 C12R001/91.
- └ 959. [WO 9002185 A](#) [AU 634171 B](#) [AU 8940478 A](#) [CA 1335965 C](#) [CN 1040823 A](#) [DE 68909797 E](#) [DK 9100223 A](#) [EP 429497 A](#) [EP 429497 B1](#) [JP 04500154 W](#) [US 5349128 A](#). Coat protein gene of cucumber mosaic virus strain WL - cloned to produce transformed plants which are resistant to CMV viral

infection. GONSALVES, D, et al. A01H001/00 A01H001/04 A01H004/00 A01H005/00 A01N063/00 C07H021/04 C12N001/21 C12N005/00 C12N005/10 C12N005/14 C12N015/00 C12N015/40 C12N015/82.

┐ 960. WO 9002184 A DE 68928445 E AU 8939870 A CN 1044126 A EP 429478 A EP 429483 A JP 04500151 W JP 04500152 W AU 634168 B EP 429478 B1 CA 1329561 C EP 693555 A1 EP 699757 A1 EP 429483 B1. Potyvirus coat protein genes - used to produce transformed plants resistant to viral infection by potyvirus and related viruses. GONSALVES, D, et al. A01H004/00 A01H005/00 A01N063/00 C07K014/08 C12N005/10 C12N005/14 C12N015/40 C12N015/82.

✓ 961. JP 02049591 A. New plant virus RNA vector - obtd. by connecting foreign gene downstream of coat protein gene of tobamo virus RNA. C12N015/83.

┐ 962. WO 8912100 A AU 8939641 A EP 444040 A FR 2631973 A. Transgenic plants resistant to potyviruses - and complete genome RNA sequence of potato virus Y. MASSON, J, et al. C12N015/00.

┐ 963. WO 8905858 A JP 3055787 B2 AU 8929276 A DK 9001265 A EP 391972 A CN 1042182 A JP 03501680 W EP 391972 B1 DE 3850261 G. Coat protein gene from C strain of cucumber mosaic virus - used to prepare plant transformation vectors and virus-resistant plants. QUEMADA, H D, et al. A01H001/00 A01H005/00 A01N001/00 C12N001/20 C12N001/21 C12N005/00 C12N005/10 C12N005/14 C12N015/00 C12N015/09 C12N015/83 C12R001/01 C12N001/21 C12R001:19.

✓ 964. WO 8904868 A AU 8928100 A CA 1337280 C. Chimeric gene construct for producing transgenic plants - contains delta-endotoxin of *Bacillus thuringiensis* for toxicity to, e.g. lepidoptera spp. BARTON, K A, et al. A01H001/04 A01H005/00 C07H015/12 C12N005/00 C12N005/10 C12N015/00 C12N015/11 C12N015/32 C12N015/84.

✓ 965. JP 01120290 A. Plant virus RNA vector - obtd. by substituting foreign gene for coat protein gene region and 30K protein gene region of tobacco mosaic virus RNA. C12N015/00 C12R001/91.

┐ 966. EP 279433 A DE 3865694 G EP 279433 B ES 2026581 T3 JP 01027476 A. DNA coding for the coat protein of cucumber mosaic virus strain Y - used for producing plants resistant to cucumber mosaic virus infection. FURUSAWA, I, et al. A01H005/00 C12N001/20 C12N005/00 C12N015/40 C12N015/82.

┐ 967. EP 278667 A US 5804439 A AU 8811383 A JP 63301787 A EP 278667 B1 DE 3850683 G ES 2060646 T3 CA 1337933 C US 5602242 A US 5627060 A. Hybrid RNA viral sequence esp. for infecting plants - comprising an infectious viral sequence and a heterologous origin of assembly and coat protein gene. AHLQUIST, P G, et al. A01H001/00 A01H005/00 C07G017/00 C07H021/02 C12N005/10 C12N007/00 C12N007/01 C12N015/00 C12N015/33 C12N015/40 C12N015/79 C12N015/82 C12N015/83.

✓ 968. JP 63014693 A. Plant virus RNA vector - prepd. by substituting coat protein gene with tobamo-virus RNA for an adventitious gene. C12N005/00 C12N007/00 C12N015/00.

┐ 969. WO 8707644 A US 5891665 A AU 8774896 A EP 270611 A GB 2199328 A JP 01500961 W GB 2199328 B EP 270611 B1 DE 3750422 G US 5489527 A US 5612193 A JP 10146197 A JP 2814433 B2. Enhancing translation of mRNA by attaching leader sequence - from RNA virus, opt. in form of complementary DNA-, e.g. for improving protein expression in transformed cells. WILSON, T M, et al.

A01H001/00 C07H021/02 C07K014/08 C12N001/20 C12N001/21 C12N005/00 C12N005/10
C12N015/00 C12N015/09 C12N015/11 C12N015/67 C12N015/70 C12N015/82 C12N015/85 C12P021/00
C12P021/02 C12P021/04 C12P021/06 C12R001/91 C12N015/09 C12R001:91 C12P021/02 C12R001:19
C12N015/09 C12R001:91.

└ 970. EP 240331 A EP 240331 B2 AU 8770934 A JP 62285791 A EP 240331 B1 DE 3751099 G ES 2068811 T3 . Virus resistant plants - contg. DNA comprising a promoter and a viral coat protein structural gene. JARVIS, N P, et al. A01H001/06 A01H005/00 C07H021/04 C12N001/20 C12N005/00 C12N005/04 C12N007/00 C12N015/00 C12N015/82.

└ 971. EP 223452 A JP 3281512 B2 AU 8664528 A ZA 8608242 A JP 62201527 A DK 8605151 A JP 06339379 A EP 223452 B1 JP 08051986 A DE 3650507 G ES 2087055 T3 IL 80433 A JP 2000041512 A IE 81098 B JP 2000157288 A . Prodn. of plants resistant to viral infection - by using recombinant DNA molecule to give genetically transformed plants. BEACHY, R N, et al. A01H001/00 A01H005/00 A01N065/00 C07H021/00 C12N001/20 C12N001/21 C12N001/22 C12N005/00 C12N005/10 C12N005/14 C12N015/00 C12N015/09 C12N015/11 C12N015/33 C12N015/63 C12N015/82 C12N015/83 C12R001:01 C12N005/10 C12R001:91 C12N001/21 C12R001:01 C12N001/20 C12R001:01 C12N001/21 C12R001:01.

└ 972. EP 221044 A AU 8664380 A JP 62175187 A ZA 8608126 A EP 221044 B1 DE 3686633 G . New plant plasmid - comprising heterologous DNA sequence and a segment of a coat protein-encoding gemini:virus DNA which permits autonomous replication. BISARO, D M, et al. A01H001/00 A01H005/00 C07G017/00 C12N005/00 C12N015/00 C12N015/83 C12P021/00.

Generate Collection

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Terms	Documents
l1 with l6 with L7	972

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09380 709
A-H86

Set Items Description

? s plant or plants

667272 PLANT
2159080 PLANTS
S1 2387728 PLANT OR PLANTS
? s virus? or viral?

1302614 VIRUS?
530308 VIRAL?
S2 1415137 VIRUS? OR VIRAL?
? s display?

S3 352757 DISPLAY?
? s insert?

S4 265542 INSERT?
? s s1 and s2 and s3

2387728 S1
1415137 S2
352757 S3
S5 1024 S1 AND S2 AND S3
? s s1 and s2 and s4

<-----User Break----->
u!
? s s1 and s2 and s4

2387728 S1
1415137 S2
265542 S4
S6 1850 S1 AND S2 AND S4
? s s5 or s6

1024 S5
1850 S6
S7 2824 S5 OR S6
? s peptide

S8 588859 PEPTIDE
? s foreign

S9 91606 FOREIGN
? s s1 and s2 and s3 and s8

2387728 S1
1415137 S2
352757 S3
588859 S8
S10 79 S1 AND S2 AND S3 AND S8
? s s1 and s2 and s3 and s9

2387728 S1
1415137 S2
352757 S3
91606 S9
S11 16 S1 AND S2 AND S3 AND S9
? s s1 and s2 and s4 and s8

2387728 S1
1415137 S2
265542 S4
588859 S8
S12 130 S1 AND S2 AND S4 AND S8
? s s1 and s2 and s4 and s0

>>>"S0" does not exist
2387728 S1
1415137 S2
265542 S4
0 S0
S13 0 S1 AND S2 AND S4 AND S0
? s s1 and s2 and s4 and s9

2387728 S1

1415137 S2
265542 S4
91606 S9
S14 197 S1 AND S2 AND S4 AND S9
? ds

Set Items Description
S1 2387728 PLANT OR PLANTS
S2 1415137 VIRUS? OR VIRAL?
S3 352757 DISPLAY?
S4 265542 INSERT?
S5 1024 S1 AND S2 AND S3
S6 1850 S1 AND S2 AND S4
S7 2824 S5 OR S6
S8 588859 PEPTIDE
S9 91606 FOREIGN
S10 79 S1 AND S2 AND S3 AND S8
S11 16 S1 AND S2 AND S3 AND S9
S12 130 S1 AND S2 AND S4 AND S8
S13 0 S1 AND S2 AND S4 AND S0
S14 197 S1 AND S2 AND S4 AND S9
? s s10 or s11 or s12 or s14

79 S10
16 S11
130 S12
197 S14
S15 376 S10 OR S11 OR S12 OR S14
? rd

...examined 50 records (50)
...examined 50 records (100)
...examined 50 records (150)
...examined 50 records (200)
...examined 50 records (250)
...examined 50 records (300)
...examined 50 records (350)
...completed examining records
S16 287 RD (unique items)
? s s16 and py<1993

Processing
287 S16
21486062 PY<1993
S17 126 S16 AND PY<1993
? t s17/3,ab/1-126

17/3,AB/1 (Item 1 from file: 5)
DIALOG(R)File 5:Biosis Previews(R)
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08756391 BIOSIS NO.: 199395045742
Tagging of %%%plant%% potyvirus replication and movement by
%%insertion%% of beta-glucuronidase into the %%%viral%%
polyprotein.
AUTHOR: Dolja Valerian V; McBride Helen J; Carrington James C
AUTHOR ADDRESS: Dep. Biol., Texas A and M University, College
Station,
Texas 77843**USA
JOURNAL: Proceedings of the National Academy of Sciences of the United
States of America 89 (21):p10208-10212 %%%1992%%
ISSN: 0027-8424
DOCUMENT TYPE: Article
RECORD TYPE: Abstract
LANGUAGE: English

ABSTRACT: Infectious RNA transcripts were generated from full-length
cDNA
clones of the tobacco etch potyvirus genome containing an
%%insertion%%
of the bacterial beta-glucuronidase (GUS) gene between the
polyprotein-coding sequences for the N-terminal 35-kDa proteinase and the
helper component-proteinase. The recombinant %%%virus%% was able
to
spread systemically in %%%plants%% and accumulated to a level
comparable

09/580704
A-H#6

1. Document ID: US 6225528 B1

Nov 14, 2000

L9: Entry 1 of 52

File: USPT

May 1, 2001

US-PAT-NO: 6225528
DOCUMENT-IDENTIFIER: US 6225528 B1
TITLE: Method of making pathogen-resistant plants by transformation with a fatty acid desaturase gene
DATE-ISSUED: May 1, 2001

US-CL-CURRENT: 800/279

APPL-NO: 9/ 143567
DATE FILED: August 28, 1998

PARENT-CASE:
This application claims priority to U.S. Provisional Application No. 60/057,510, filed Sep. 4, 1997, which is incorporated by reference herein.

IN: Chin; Chee-Kok, Wang; Chunlin, Xing; Jinsong

AB: The present invention provides pathogen-resistant transgenic plants and methods of making the plants. The transgenic plants display enhanced resistance to a variety of fungal, bacterial and viral plant pathogens due to expression of a gene that increases the unsaturated fatty acid content of the plant's cells, as compared with an equivalent, but non-transformed plant. The preferred embodiment of the invention is a plant expressing a heterologous Δ 9 desaturase gene from yeast, which particularly increases cytosolic quantities of 16:1, 16:2 and 18:1 fatty acids.

L9: Entry 1 of 52

File: USPT

May 1, 2001

DOCUMENT-IDENTIFIER: US 6225528 B1
TITLE: Method of making pathogen-resistant plants by transformation with a fatty acid desaturase gene

ABPL:
The present invention provides pathogen-resistant transgenic plants and methods of making the plants. The transgenic plants display enhanced resistance to a variety of fungal, bacterial and viral plant pathogens due to expression of a gene that increases the unsaturated fatty acid content of the plant's cells, as compared with an equivalent, but non-transformed plant. The preferred embodiment of the invention is a plant expressing a heterologous Δ 9 desaturase gene from yeast, which particularly increases cytosolic quantities of 16:1, 16:2 and 18:1 fatty acids.

2. Document ID: US 6147278 A

L9: Entry 2 of 52

File: USPT

US-PAT-NO: 6147278
DOCUMENT-IDENTIFIER: US 6147278 A
TITLE: Plant vectors
DATE-ISSUED: November 14, 2000

US-CL-CURRENT: 800/278; 435/320.1, 435/468, 435/469, 435/69.1, 536/23.72, 800/288

APPL-NO: 9/ 261770
DATE FILED: March 3, 1999

PARENT-CASE:
CROSS-REFERENCE TO RELATED APPLICATION This application is a continuation of application Ser. No. 07/711,576 filed May 31, 1991, now abandoned, which is a continuation of application Ser. No. 07/209,239 filed Jun. 26, 1988, now abandoned, which is a continuation-in-part of application Ser. No. 06/899,270 filed Aug. 26, 1986, now abandoned, which is a continuation-in-part of application Ser. No. 06/791,249 filed Oct. 25, 1985, now abandoned.

IN: Rogers; Stephen G., Brand; Leslie, Horsch; Robert B., Fraley; Robert T., Elmer; James Scott, Bisaro; David

AB: The invention relates to novel plant plasmid vectors comprising geminivirus DNA or a portion thereof having inserted therein a heterologous DNA sequence or gene, to processes and DNA intermediates useful in producing said vectors and to methods utilizing such vectors to replicate and express heterologous DNA sequences or genes in plants. In some embodiments, methods and compositions are provided for Ti plasmid delivery of these novel vectors into plants. In other embodiments, methods and compositions are provided which allow for the generation of geminivirus DNA containing plant plasmids in stably transformed plants. In still other embodiments, methods and compositions are provided for replicating and expressing heterologous DNA sequences or genes in plants employing the geminivirus DNA containing vectors of the present invention without causing disease symptoms.

L9: Entry 2 of 52

File: USPT

Nov 14, 2000

DOCUMENT-IDENTIFIER: US 6147278 A
TITLE: Plant vectors

DEPR:
While neither the transgenic A- or B-containing plants exhibited virus disease symptoms, it was demonstrated, in Example 14, supra, that inoculation of B-containing plants with vectors comprising the TGMV A-component subsequently displayed virus symptoms. Subsequent experiments by Sunter et al (1987) have shown that one-quarter of the progeny produced by crossing a transgenic A plant with a transgenic B plant show geminivirus symptoms and contain infectious virus particles. These results show that the integrated tandem copies of the TGMV DNA's are functional, are able to be released from their integrated state and maintain their ability to produce infectious virus when genetically combined in the same cell. These results further demonstrate that the A component contains the necessary sequences and/or genes to

enable release of B component DNA from its integrated state. Specifically, the ability of TGMV-A component DNA (e.g. coat protein-encoding geminivirus DNA molecules) to cause the release and subsequent replication of TGMV-B component DNA molecules, demonstrates that a geminivirus trans-acting factor(s) is (are) encoded in the TGMV-A component DNA. Additionally, these results demonstrate that both the TGMV-A and -B component DNA's contain a sequence or sequences responsive to the geminivirus trans-acting factor(s). A geminivirus DNA sequence responsive to a geminivirus trans-acting factor is understood herein to mean a geminivirus DNA sequence, the presence of which coupled to the presence of a geminivirus trans-acting factor, results in the autonomous replication of a DNA molecule containing the responsive sequence. It was further demonstrated, in Examples 13 and 14 supra, that heterologous DNA sequences can be inserted into and/or in place of the coat protein gene without disrupting the ability of the TGMV A-component to form plasmid DNA molecules in plant cells or cause disease symptoms in transgenic plants containing tandem copies of the TGMV B-component. These results demonstrate that the DNA sequences coding for the geminivirus coat protein per se are not required for replication of geminivirus DNA or geminivirus-containing plasmid DNA molecules in plants and/or plant cells. Specifically, Examples 13 and 14, supra, teach that neither the geminivirus trans-acting factor nor sequences responsive to said factor are contained within the DNA sequences encoding the TGMV coat protein. These foregoing examples further set forth a method by which one of skill in the art can determine the minimal sequence or sequences required for binary geminivirus replication in a plant cell. Specifically, the foregoing examples demonstrate that by performing conventional deletion and/or mutation analysis, a minimal binary geminivirus replicon can be determined.

3. Document ID: US 6146628 A

L9: Entry 3 of 52

File: USPT

Nov 14, 2000

US-PAT-NO: 6146628

DOCUMENT-IDENTIFIER: US 6146628 A

TITLE: Biotherapeutic agents comprising recombinant PAP and PAP mutants

DATE-ISSUED: November 14, 2000

US-CL-CURRENT: 424/134.1; 424/142.1, 424/143.1, 424/147.1, 424/148.1, 424/183.1, 424/184.1, 424/187.1

APPL-NO: 8/ 501253

DATE FILED: July 11, 1995

IN: Uckun; Fatih M., Turner; Nilgun E.

AB: Biotherapeutic agents are provided which comprise recombinant PAP or a biologically equivalent variant or mutant thereof, linked to a targeting moiety which are effective for the treatment of certain human diseases. The invention further provides a

process for producing the biotherapeutic agents as well as a method which utilizes the disclosed biotherapeutic agents to systemically treat cancer patients.

L9: Entry 3 of 52

File: USPT

Nov 14, 2000

DOCUMENT-IDENTIFIER: US 6146628 A

TITLE: Biotherapeutic agents comprising recombinant PAP and PAP mutants

BSPR:

PAP displays broad-spectrum antiviral activity against plant viruses, herpes simplex virus, cytomegalovirus, poliovirus, and influenza virus. Aron et al., Agents Chemotherapv, 17, 1032 (1980). In fact, pokeweed antiviral protein was discovered due to its ability to inhibit the transmission of tobacco mosaic virus (TMV) in plants and it was subsequently demonstrated that the purified protein was equally effective against a number of other plant viruses. Tomlinson et al., J. Gen. Virol., 22, 225 (1974). All of these experiments were performed in a similar manner; PAP was mixed with the virus inoculum which was then rubbed on plant leaves in the presence of an abrasive substance, such as carborundum, which damages the tissue allowing the entry of the virus and presumably the PAP. Using this method, it was found that highly diluted solutions of PAP were capable of inhibiting local lesion formation caused by southern bean mosaic virus as well as cucumber mosaic virus. Wyatt et al., Phytopath., 59, 1787 (1969); Tomlinson et al., cited supra. Using the local lesion assay system on Phaseolus vulgaris, it has been shown that PAP inhibited viral infection at very low concentrations. Irvin et al., Arch. Biochem. Biophys., 200, 418 (1980). PAP has also been shown to effectively inhibit TWV infection of tobacco protoplasts with nearly complete inhibition obtained with 10 .mu.g/mL (.apprxq.300 nM). Grasso et al., Phytopath., 98, 53 (1980). Furthermore, in a study done to compare the relative antiviral properties of a number of ribosome inactivating proteins (RIPs) including PAP upon the formation of local lesions on Nicotiana glutinosa by TMV, it was found that all of the RIPs tested had antiviral activity, but none of the studied RIP's were as effective as PAP. Stevens et al., Experientia, 37, 257 (1981).

4. Document ID: US 6133505 A

L9: Entry 4 of 52

File: USPT

Oct 17, 2000

US-PAT-NO: 6133505

DOCUMENT-IDENTIFIER: US 6133505 A

TITLE: Phytopathogenic geminivirus resistant transgenic plants and seeds and methods for

obtaining same by introduction of mutated CI gene
DATE-ISSUED: October 17, 2000

US-CL-CURRENT: 800/280; 435/440

APPL-NO: 8/ 809103
DATE FILED: March 17, 1997

FOREIGN-APPL-PRIORITY-DATA:
COUNTRY

	APPL-NO	APPL-DATE
FR	94 11040	September 15, 1994

PCT-DATA:
APPL-NO
DATE-FILED
PUB-NO
PUB-DATE
371-DATE
102(E)-DATE

PCT/FR95/01192	September 15, 1995	WO96/08573	Mar 21, 1996	Mar 17, 1997	Mar 17, 1997
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IN: Gronenborn; Bruno

AB: Nucleotide sequences produced by mutation (also known as mutant nucleotide sequences) of C1 nucleotide sequences present in a pathogenic geminivirus genome in plants with one or more mutations capable of producing a dominant negative phenotype for the replication of the pathogenic virus, its diffusion in a plant, or its spread from one plant to another, especially through vectors such as insects, the mutant nucleotide sequences being capable of fully or partially inhibiting the replication and/or diffusion and/or spread of the pathogenic virus for producing phytopathogenic geminivirus resistant or tolerant transgenic plants.

L9: Entry 4 of 52
File: USPT
Oct 17, 2000

DOCUMENT-IDENTIFIER: US 6133505 A
TITLE: Phytopathogenic geminivirus resistant transgenic plants and seeds and methods for obtaining same by introduction of mutated C1 gene

DEPR:
To confirm these results, leaf discs of potentially resistant plants are agroinoculated with the STYLCV. Seven days after agroinoculation, the various forms of viral DNA (ss and ds) are detected in the leaf discs of non-transgenic *N. benthamiana*. This provides evidence of good infectivity of the clone used for inoculating the leaf discs. Three plants display very reduced replication relative to a positive control: these are resistant plants (absence of symptoms, but residual replication of the viral DNA). Two plants do not have any form of viral DNA: these are completely resistant plants. Curiously, two plants have the various forms of viral DNA, whereas no signal was visible in squash or in Southern blot. This might be explained by the "inoculum pressure" used in the experiments of agroinoculation of leaf discs. This "inoculum pressure" is higher than was used in the agroinoculation of whole plants, and so would lead to a

change of the ratio of mutated to wild-type C1 proteins in favour of the wild-type protein, and consequently to replication of the viral DNA. On the other hand this (ambiguous) behaviour might correspond to an inhibition of the movement (at long range?) of the virus in planta, since the agroinoculation of leaf discs makes it possible to avoid phenomena of movement at long range.

5. Document ID: US 6110466 A

L9: Entry 5 of 52
File: USPT
Aug 29, 2000

US-PAT-NO: 6110466
DOCUMENT-IDENTIFIER: US 6110466 A
TITLE: Modified plant viruses as vectors
DATE-ISSUED: August 29, 2000

US-CL-CURRENT: 424/199.1; 424/185.1, 424/186.1, 424/188.1, 424/192.1, 424/202.1, 424/204.1, 435/235.1, 435/69.1

APPL-NO: 8/ 137032
DATE FILED: December 15, 1993

FOREIGN-APPL-PRIORITY-DATA:
COUNTRY

	APPL-NO	APPL-DATE
GB	9108386	April 19, 1991

PCT-DATA:
APPL-NO
DATE-FILED
PUB-NO
PUB-DATE
371-DATE
102(E)-DATE

PCT/GB92/00589	April 2, 1992	WO92/18618	Oct 29, 1992	Dec 15, 1993	Dec 15, 1993
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IN: Lomonosoff; George Peter, Johnson; John Emil

AB: The invention relates to assembled particles of a plant virus containing a predetermined foreign peptide as part of the coat protein of the virus, and a method for their production. The foreign peptide is preferably a biologically functional peptide, the biological application of which requires or is enhanced by presentation of the peptide in association with a larger molecule or particle.

L9: Entry 5 of 52
File: USPT
Aug 29, 2000

DOCUMENT-IDENTIFIER: US 6110466 A
TITLE: Modified plant viruses as vectors

ABPR:

The invention relates to assembled particles of a plant virus containing a predetermined foreign peptide as part of the coat protein of the virus, and a method for their production. The foreign peptide is preferably a biologically functional peptide, the biological application of which requires or is enhanced by presentation of the peptide in association with a larger molecule or particle.

BSPR:

This invention relates to the use of viruses as carriers (vectors) for the production or presentation of foreign peptides. More particularly, the invention relates to the genetic manipulation of viral nucleic acid by incorporation of foreign nucleic acid sequences which are expressed as peptides in the virus particle (virion). In this specification the term "foreign", as applied to a peptide or to the nucleic acid encoding therefor, signifies peptide or nucleic acid sequences which are not native to the plant virus used as a vector. Such sequences can be alternatively described as exogenous or heterologous sequences. The term "peptide" includes small peptides and polypeptides.

BSPR:

The present invention utilises plant viruses as vector systems for the expression of foreign nucleotide sequences ie nucleotide sequences (RNA or DNA) which are not present in plant viruses, as found in Nature, and which in consequence code for peptides not normally found in any naturally occurring plant virus.

BSPR:

The present invention comprises assembled particles of a plant virus containing a foreign peptide. The plant viruses of the present invention are therefore modified forms of the native viruses and for convenience will be referred to as modified viruses.

BSPR:

The foreign peptides which may be incorporated into plant viruses according to this invention may be of highly diverse types and are subject only to the limitation that the nature and size of the foreign peptide and the site at which it is placed in or on the virus particle do not interfere with the capacity of the modified virus to assemble when cultured in vitro or in vivo. In broad concept, modified viruses may be formed from any biologically useful peptides (usually polypeptides) the function of which requires a particular conformation for its activity. This may be achieved by association of the peptide with a larger molecule eg to improve its stability or mode of presentation in a particular biological system. Examples of such peptides are peptide hormones; enzymes; growth factors; antigens of protozoal, viral, bacterial, or fungal origin; antibodies including anti-idiotypic antibodies; immunoregulators and cytokines eg interferons and interleukins; receptors; adhesins; and parts or precursors of any of the foregoing types of peptide. The peptide preferably contains more than 5 amino acids.

BSPR:

To produce modified plant virus particles in accordance with this invention the plant viral nucleic acid is modified by introducing a nucleotide sequence coding for the foreign peptide eg

an animal virus antigen at that part of the plant viral genome which codes for an exposed portion of the coat protein, infecting plants or plant cells with the modified viral nucleic acid, and harvesting assembled particles of the modified virus. This procedure is best carried out by direct manipulation of the DNA of the virus in the case of DNA viruses or by manipulation of a cDNA corresponding to the RNA of an RNA virus. In the case of an RNA virus, an RNA transcript of the modified DNA is usually prepared for inoculation of plant cells or preferably whole plants so as to achieve a multiplication stage prior to the harvesting of assembled particles of the modified virus. In the case of a DNA virus, the DNA itself is introduced into the plant. In this way, the foreign peptide is initially expressed as part of the capsid protein and is thereby produced as part of the whole virus particle. The peptide may thus be produced as a conjugate molecule intended for use as such. Alternatively, the genetic modification of the virus may be designed in order to permit release of the desired peptide by the application of appropriate agents which will effect cleavage from the virus particle.

CLPR:

1. Assembled particles of a plant virus containing a foreign peptide encoded by an exogenous nucleotide sequence as part of the coat protein of the virus, the particles having been assembled in whole plants or in plant cells, and wherein the coat protein of the virus has a .beta.-barrel structure and said virus is selected from the group consisting of Comoviruses, Tombusviruses, Sobemoviruses, and Nepoviruses, and the site of insertion of the foreign peptide is in a loop between individual strands of .beta. sheet.

CLPR:

8. Virus particles according to claim 1, in which the foreign peptide is inserted in the .beta.B- .beta.C loop of the plant virus.

6. Document ID: US PP11418 P

L9: Entry 6 of 52

File: USPT

Jun 13, 2000

US-PAT-NO: PP11418
DOCUMENT-IDENTIFIER: US PP11418 P
TITLE: Raspberry plant named 'Glen Ample'
DATE-ISSUED: June 13, 2000

US-CL-CURRENT: PLT/204

APPL-NO: 9/ 069762
DATE FILED: April 30, 1998

IN: McNicol; Ronnie J., Jennings; Derek L.

AB: The new and distinct cultivar of raspberry (i.e., *Rubus idaeus* L.) is provided.

The cultivar forms attractive large bright red berries of good flavor in exceptionally high yields on long fruiting laterals. The drupelet cohesion tends to be somewhat reduced when the plant is grown in cooler climates (e.g., Scotland). The plant exhibits a spine-free very

upright growth habit of good vigor. The berries are suitable for consumption as high grade fresh fruit and also are amenable to processing. Additionally, the plant has displayed resistance to *Amphorophora idaei* aphid virus vector.

L9: Entry 6 of 52

File: USPT

Jun 13, 2000

DOCUMENT-IDENTIFIER: US PP11418 P
TITLE: Raspberry plant named 'Glen Ample'

ABPL:

The new and distinct cultivar of raspberry (i.e., *Rubus idaeus* L.) is provided. The cultivar forms attractive large bright red berries of good flavor in exceptionally high yields on long fruiting laterals. The drupelet cohesion tends to be somewhat reduced when the plant is grown in cooler climates (e.g., Scotland). The plant exhibits a spine-free very upright growth habit of good vigor. The berries are suitable for consumption as high grade fresh fruit and also are amenable to processing. Additionally, the plant has displayed resistance to *Amphorophora idaei* aphid virus vector.

7. Document ID: US 6057492 A

L9: Entry 7 of 52

File: USPT

May 2, 2000

US-PAT-NO: 6057492
DOCUMENT-IDENTIFIER: US 6057492 A
TITLE: Plants resistant to tospoviruses
DATE-ISSUED: May 2, 2000

US-CL-CURRENT: 800/280; 435/320.1, 435/419, 435/468, 435/69.1, 536/23.72, 800/265, 800/279, 800/301

APPL-NO: 8/ 913374
DATE FILED: September 18, 1997

FOREIGN-APPL-PRIORITY-DATA:
COUNTRY

	APPL-NO	APPL-DATE
GB	9505907	March 23, 1995

PCT-DATA:
APPL-NO

	DATE-FILED	PUB-NO	PUB-DATE	371-DATE	102(E)-DATE
PCT/EP96/01271	March 22, 1996				
		WO96/29420	Sep 26, 1996		
			Sep 18, 1997		
				Sep 18, 1997	

IN: de Haan; Petrus Theodorus

AB: Plant transformation vectors comprising a polynucleotide effective to render resistance or tolerance to infection by a tospovirus, and a microbiological process for making virus tolerant or resistant plants are provided herein.

L9: Entry 7 of 52

File: USPT

May 2, 2000

DOCUMENT-IDENTIFIER: US 6057492 A
TITLE: Plants resistant to tospoviruses

CLPR:

17. A microbiological process for making virus tolerant or resistant plants, comprising the steps of: (i) transforming regenerable plant material with a polynucleotide comprising a sequence which hybridizes under stringent conditions to a polynucleotide having the sequence of SEQ ID NO.: 1, or to a polynucleotide comprising nucleotides 3813 to 4721 of SEQ ID NO.: 1 (ii) regenerating the transformed regenerable plant material into a morphologically normal fertile plant; and (iii) selecting a plant that displays resistance or tolerance when exposed to a virus.

8. Document ID: US 6013864 A

L9: Entry 8 of 52

File: USPT

Jan 11, 2000

US-PAT-NO: 6013864
DOCUMENT-IDENTIFIER: US 6013864 A
TITLE: Plants resistant to infection by luteoviruses
DATE-ISSUED: January 11, 2000

US-CL-CURRENT: 800/301; 435/320.1, 435/419, 435/468, 536/23.72, 800/280, 800/288, 800/317.2, 800/317.3, 800/317.4, 800/320, 800/320.2, 800/320.3

APPL-NO: 8/ 617454
DATE FILED: March 18, 1996

PARENT-CASE:

The present application is a continuation-in-part of U.S. patent application Ser. No. 08/326,297, filed Oct. 20, 1994, now U.S. Pat. No. 5,510,253, which is a continuation of U.S. patent application Ser. No. 08/012,688, filed Feb. 3, 1993, now abandoned.

IN: Mitsky; Timothy Albert, Hemenway; Cynthia Lou, Tumer; Nilgun Erenen, Lawson; Edgar Clifford

AB: An isolated DNA sequence which codes for a luteo replicase gene is disclosed herein. A method for providing resistance to infection by a virus by expressing a replicase gene in plants is also disclosed, as are transgenic plants containing the replicase gene.

L9: Entry 8 of 52

File: USPT

Jan 11, 2000

DOCUMENT-IDENTIFIER: US 6013864 A
TITLE: Plants resistant to infection by luteoviruses

DEPR:
PLRV is not mechanically transmissible. Spread of PLRV from infected plants to uninfected plants can only be accomplished by aphids. This is the reason that insecticide application is currently necessary for controlling this disease. Virus resistant potatoes will no longer require insecticides to control aphids. Potato cultivars which are resistant to PLRV often display no or reduced titers of PLRV and the virus is not as easily transmitted from these plants to other plants by aphids. This characteristic is of commercial significance because it limits the potential for virus epidemics in the field.

9. Document ID: US 6001986 A

L9: Entry 9 of 52

File: USPT

Dec 14, 1999

US-PAT-NO: 6001986
DOCUMENT-IDENTIFIER: US 6001986 A
TITLE: Antiviral proteins, amarandin 1 and 2, from *Amaranthus viridis*, DNAs encoding therefrom
DATE-ISSUED: December 14, 1999

US-CL-CURRENT: 536/23.6; 435/252.3, 435/320.1, 435/471

APPL-NO: 8/ 916443
DATE FILED: August 22, 1997

IN: Kim; Yong Sig, Park; Sun Chung, Oh; Soo Kyung, Lee; Hosull, Cho; Jeong Woo, Chung; Chang H.

AB: DNA sequences encoding antiviral proteins, amarandin 1 and 2 from *Amaranthus viridis* is disclosed. Expression vectors encoding amarandin 1 or 2 and transformed host cells are also disclosed.

L9: Entry 9 of 52

File: USPT

Dec 14, 1999

DOCUMENT-IDENTIFIER: US 6001986 A
TITLE: Antiviral proteins, amarandin 1 and 2, from *Amaranthus viridis*, DNAs encoding therefrom

BSPR:
During our screening studies searching for new RIP we found that *Amaranthus viridis* crude extracts displayed both translational inhibitory and antiviral activities against plant viruses.

10. Document ID: US 5998699 A

L9: Entry 10 of 52

File: USPT

Dec 7, 1999

US-PAT-NO: 5998699
DOCUMENT-IDENTIFIER: US 5998699 A
TITLE: Potyvirus coat protein genes and plants transformed therewith
DATE-ISSUED: December 7, 1999

US-CL-CURRENT: 800/301; 435/419, 536/23.72, 800/280

APPL-NO: 8/ 358653
DATE FILED: December 19, 1994

PARENT-CASE:
The present application is a continuation of U.S. Ser. No. 08/232,846, filed Apr. 25, 1994, now abandoned which is a continuation of U.S. Ser. No. 08/013,971, filed Feb. 4, 1993, now abandoned which is a continuation of U.S. Ser. No. 07/656,167, filed Feb. 19, 1991, now abandoned, which is a continuation of international application PCT/US89/03094, filed Jul. 20, 1989, which is a continuation of U.S. Ser. No. 07/368,710, filed Jun. 19, 1989, now abandoned, which is a continuation in part of U.S. Ser. No. 07/234,412, filed Aug. 19, 1988, and a continuation of U.S. Ser. No. 07/323,536, filed Mar. 14, 1989, each now abandoned.

IN: Slightom; Jerry L., Quemada; Hector D., Gonsalves; Dennis, Lhostis; Brigitte

AB: The present invention relates to the coat protein genes of Papaya Ringspot Virus Strain papaya ringspot (PRV-p), Watermelon Mosaic Virus II (WMVII), and Zucchini Yellow Mosaic Virus (ZYMV); to expression vectors which contain a coat protein gene for PVP-p, WMVII or ZYMV, and, additionally, the necessary genetic regulatory sequences needed for expression of a gene transferred into a plant; to bacterial or plant cells which are transformed with an expression vector containing the PVP-p, WMVII or ZYMV coat protein genes; to transgenic plants which are produced from plant cells transformed with an expression vector containing the coat protein gene from PVP-p, WMVII or ZYMV; and to a process of producing transgenic plants which have increased resistance to viral infection.

L9: Entry 10 of 52

File: USPT

Dec 7, 1999

DOCUMENT-IDENTIFIER: US 5998699 A
TITLE: Potyvirus coat protein genes and plants transformed therewith

BSPR:
Tumer et al. (1987) "Expression of alfalfa mosaic virus coat protein gene confers cross-protection in transgenic tobacco and tomato plants", EMBO J. 6:1181-1188, disclose transgenic tobacco and tomato plants transformed with the coat protein gene of alfalfa mosaic virus displayed increased resistance to infection by alfalfa mosaic virus.

11. Document ID: US 5990388 A

L9: Entry 11 of 52

File: USPT

Nov 23, 1999

US-PAT-NO: 5990388

DOCUMENT-IDENTIFIER: US 5990388 A

TITLE: Resistance to viruses and viroids in transgenic plants and animals expressing

dsRNA-binding protein

DATE-ISSUED: November 23, 1999

US-CL-CURRENT: 800/301; 435/320.1, 800/280, 800/317.2, 800/317.3

APPL-NO: 8/ 482286

DATE FILED: June 7, 1995

IN: Roth; Don Allen, Langland; Jeffrey Olaf

AB: The invention provides a method for imparting resistance in animals to viruses, and in plants to viruses and viroids, that express double-stranded RNA-like structures (dsRNAs). This method enables the binding of pathogenic dsRNAs during the infection process by expression of dsRNA-binding protein in transgenic animal and plant hosts, thus interrupting the infection cycle and inhibiting disease. The presence of a dsRNA-binding protein in a transgenic host renders the transgenic host resistant to the phenotypic symptoms of viral infection and/or decreased pathogen replication. Accordingly, the present invention provides a genetically engineered animal and plant, stably transformed to express a dsRNA-binding protein, such that the transgenic host displays resistance to virus and/or viroid challenge.

L9: Entry 11 of 52

File: USPT

Nov 23, 1999

DOCUMENT-IDENTIFIER: US 5990388 A

TITLE: Resistance to viruses and viroids in transgenic plants and animals expressing dsRNA-binding protein

ABPL:

The invention provides a method for imparting resistance in animals to viruses, and in plants to viruses and viroids, that express double-stranded RNA-like structures (dsRNAs). This method enables the binding of pathogenic dsRNAs during the infection process by expression of dsRNA-binding protein in transgenic animal and plant hosts, thus interrupting the infection cycle and inhibiting disease. The presence of a dsRNA-binding protein in a transgenic host renders the transgenic host resistant to the phenotypic symptoms of viral infection and/or decreased pathogen replication. Accordingly, the present invention provides a genetically engineered animal and plant, stably transformed to express a dsRNA-binding protein, such that the transgenic host displays resistance to virus and/or viroid challenge.

BSPR:

Accordingly, the present invention provides a genetically engineered animal and plant, stably transformed to express constitutively a dsRNA-binding protein, such that the transgenic host displays resistance to virus and/or viroid challenge. One aspect of the invention contemplates broad spectrum resistance in transgenic plant cells to plant viruses having a dsRNA-like structure, including but not limited to the phytoeovirus group, the tymovirus group, luteovirus group, tombusvirus group, Southern bean mosaic virus group, tobacco necrosis virus group, maize chlorotic dwarf virus group, closterovirus group, carlavirus group, potyvirus group, potexvirus group, tobamovirus group, nepovirus group, pea enation mosaic virus group, comovirus group, tobravirus group, cucumovirus group, bromovirus group, ilarvirus group, alfalfa mosaic virus group, and hordeivirus group; and to plant viroids having a dsRNA-like structure, including but not limited to the potato spindle tuber viroid, the coconut cadang-cadang viroid group, avocado sunblotch viroid group, and hop latent viroid group.

12. Document ID: US 5977438 A

L9: Entry 12 of 52

File: USPT

Nov 2, 1999

US-PAT-NO: 5977438

DOCUMENT-IDENTIFIER: US 5977438 A

TITLE: Production of peptides in plants as viral coat protein fusions

DATE-ISSUED: November 2, 1999

US-CL-CURRENT: 800/288; 435/235.1, 435/419, 435/468, 435/69.7, 435/70.1, 536/23.4, 536/23.5, 536/23.72, 800/278, 800/298

APPL-NO: 8/ 324003

DATE FILED: October 14, 1994

PARENT-CASE:

CROSS-REFERENCE TO RELATED APPLICATIONS The present application is a continuation-in-part of application Ser. No. 08/176,414, filed on Dec. 29, 1993, now U.S. Pat. No. 5,811,653, which is a continuation-in-part of application Ser. No. 07/997,733, filed Dec. 30, 1992, now abandoned. The present application is also a continuation-in-part of application Ser. No. 08/184,237, filed Jan. 19, 1994, now U.S. Pat. No. 5,589,367 which is a continuation-in-part of application Ser. No. 07/997,733, filed Dec. 30, 1992, now abandoned, which is a continuation of application Ser. No. 07/923,692, filed Jul. 31, 1992, now U.S. Pat. No. 5,316,931, which is a continuation-in-part of applications Ser. No. 07/600,244, filed Oct. 22, 1990, now abandoned, Ser. No. 07/641,617, filed Jan. 16, 1991, now abandoned, application Ser. No. 07/737,899, filed Jul. 26, 1991, now abandoned, and application Ser. No. 07/739,143, filed Aug. 1, 1991, now abandoned. Application Ser. No. 07/600,244 is a continuation of application Ser. No. 07/310,881, filed Feb. 17, 1989, now abandoned, which is a continuation-in-part of applications Ser. No. 07/160,766 and Ser. No. 07/160,771, both filed on Feb. 26, 1988 and now abandoned. Application Ser. No. 07/641,617 is a continuation of application Ser. No. 07/347,637, filed May 5, 1989, now

abandoned. Application

Ser. No. 07/737,899 is a continuation of application Ser. No. 07/363,138, filed Jun. 8, 1989, now

abandoned, which is a continuation-in-part of application Ser. No. 07/219,279, filed Jul. 15,

1988, now abandoned. Application Ser. No. 07/739,143 is a continuation-in-part of applications

Ser. No. 07/600,244, filed Oct. 22, 1990, now abandoned, Ser. No. 07/641,617, filed Jan. 16,

1991, now abandoned, and Ser. No. 07/737,899, filed Jul. 26, 1991, now abandoned.

IN: Turpen; Thomas H., Reinl; Stephen J., Grill; Laurence K.

AB: The present invention relates to foreign peptide sequences fused to recombinant

plant viral structural proteins and a method of their production. Fusion proteins are

economically synthesized in plants at high levels by biologically contained tobamoviruses.

The fusion proteins of the invention have many uses. Such uses include use as antigens for

inducing the production of antibodies having desired binding properties, e.g., protective

antibodies, or for use as vaccine antigens for the induction of protective immunity,

including immunity against parasitic infections.

L9: Entry 12 of 52

File: USPT

Nov 2, 1999

DOCUMENT-IDENTIFIER: US 5977438 A

TITLE: Production of peptides in plants as viral coat protein fusions

ABPL:

The present invention relates to foreign peptide sequences fused to recombinant plant viral

structural proteins and a method of their production. Fusion proteins are economically

synthesized in plants at high levels by biologically contained tobamoviruses. The fusion proteins

of the invention have many uses. Such uses include use as antigens for inducing the production of

antibodies having desired binding properties, e.g., protective antibodies, or for use as vaccine

antigens for the induction of protective immunity, including immunity against parasitic infections.

13. Document ID: US 5958422 A

L9: Entry 13 of 52

File: USPT

Sep 28, 1999

US-PAT-NO: 5958422

DOCUMENT-IDENTIFIER: US 5958422 A

TITLE: Modified plant viruses as vectors of heterologous peptides

DATE-ISSUED: September 28, 1999

US-CL-CURRENT: 424/199.1; 435/320.1, 435/419, 435/421, 514/2, 536/23.4, 536/23.6

APPL-NO: 8/ 612858

DATE FILED: June 5, 1996

FOREIGN-APPL-PRIORITY-DATA:

COUNTRY

APPL-NO

APPL-DATE

GB

9414118

July 13, 1994

PCT-DATA:

APPL-NO

DATE-FILED

PUB-NO

PUB-DATE

371-DATE

102(E)-DATE

PCT/GB95/01618

July 10, 1995

WO96/02649

Feb 1, 1996

Jun 5, 1996

Jun 5, 1996

IN: Lomonossoff; George Peter

AB: The invention relates to assembled particles of a plant virus containing a

foreign peptide insert in the coat protein of the virus. The site of the insert is free from

direct sequence repeats flanking the insert. The invention also relates to a method of

production of the particles and their use, in particular in vaccines.

L9: Entry 13 of 52

File: USPT

Sep 28, 1999

DOCUMENT-IDENTIFIER: US 5958422 A

TITLE: Modified plant viruses as vectors of heterologous peptides

ABPR:

The invention relates to assembled particles of a plant virus containing a foreign peptide insert

in the coat protein of the virus. The site of the insert is free from direct sequence repeats

flanking the insert. The invention also relates to a method of production of the particles and

their use, in particular in vaccines.

BSPR:

This invention relates to the use of viruses as carriers (vectors) for the production or

presentation of foreign peptides. More particularly, the invention relates to the genetic

manipulation of viral nucleic acid by incorporation of foreign nucleic acid sequences which are

expressed as peptides in the virus particle (virion). In this specification the term "foreign",

as applied to a peptide or to the nucleic acid encoding therefor, signifies peptide or nucleic

acid sequences which are not native to the plant virus used as a vector. Such sequences can be

alternatively described as exogenous or heterologous sequences. The term "peptide" includes small

peptides and polypeptides.

BSPR:

Our patent application WO 92/18618 describes the utilization of plant viruses as vector systems

for the expression of foreign nucleotide sequences, ie nucleotide sequences (RNA or DNA) which

are not present in plant viruses, as found in Nature, and which in consequence code for peptides

not normally found in any naturally occurring plant virus. The invention described therein comprises assembled particles of a plant virus containing a foreign peptide. The plant viruses of the invention are therefore modified forms of the native viruses and for convenience will be referred to as modified viruses.

BSPR:

The foreign peptides which may be incorporated into plant viruses according to our prior application WO92/18618 may be of highly diverse types and are subject only to the limitation that the nature and size of the foreign peptide and the site at which it is placed in or on the virus particle do not interfere with the capacity of the modified virus to assemble when cultured in vitro or in vivo. In broad concept, modified viruses may be formed from any biologically useful peptides (usually polypeptides) the function of which requires a particular conformation for its activity. This may be achieved by association of the peptide with a larger molecule eg to improve its stability or mode of presentation in a particular biological system. Examples of such peptides are peptide hormones; enzymes; growth factors; antigens of protozoal, viral, bacterial, fungal or animal origin; antibodies including anti-idiotypic antibodies; immunoregulators and cytokines, eg interferons and interleukins; receptors; adhesions; and parts of precursors of any of the foregoing types of peptide. The peptide preferably contains more than 5 amino acids.

BSPR:

Firstly, the process used for modifying the plant viral nucleic acid by introducing a nucleotide sequence coding for a foreign peptide should avoid the presence of direct sequence repeats flanking the insert. In the context of the present invention a construct containing a direct sequence repeat is one in which an identical oligonucleotide sequence is present on both sides of the inserted nucleotide. Such constructs can be genetically unstable because recombination can occur between the sequence repeats leading to loss of the foreign peptide coding sequence and reversion to the wild type sequence. Secondly, where the foreign oligonucleotide sequence is inserted into the plant virus genome as a substitution for part of the existing sequence, the resultant modified viral coat protein may be missing in an amino acid sequence which is important for virus replication, encapsidation and spread in the plant. This defect may be readily determined and avoided. Thirdly, the heterologous sequence should not be inserted at a sub-optimal site.

BSPR:

The present invention comprises assembled particles of a plant virus containing a foreign peptide in which the corresponding foreign nucleic acid has been inserted into the plant virus genome in the absence of direct sequence repeats flanking the insert and preferably as an addition to the existing nucleic acid.

BSPR:

In a further aspect of the present invention, cDNA clones of CPMV RNAs M and B have been constructed, in which the cDNA clone of the M RNA contains an inserted oligonucleotide sequence encoding a foreign peptide, which make use of the cauliflower mosaic virus (CaMV) 35S promoter sequence linked to the 5' ends of the viral cDNAs to generate infectious transcripts in the plant. This technique overcomes some of the problems encountered with

the use of transcripts generated in vitro and is applicable to all plant RNA viruses.

DEPR:

Oligonucleotide sequences encoding various foreign peptides (see Table 1) were substituted for the sequence between the NheI and AatII sites of pCP2-AatII as described in Example 1. The pCP2-AatII variants and pCP-1 were linearised and inoculated onto the primary leaves of cowpea plants. In all cases infections developed and stable chimaeric virus particles expressing the appropriate foreign peptide were recovered from plants.

CLPR:

1. Assembled particles of a plant virus containing a foreign peptide insert as an addition at a non-terminal site in the coat protein of the virus, the site of the insert in the coat protein corresponding to a site in the plant virus genome which is free from direct nucleotide sequence repeats flanking the insert and wherein the coat protein of the virus has a .beta.-barrel structure and the site of insertion of the foreign peptide is in a loop connecting .beta. sheets of the plant virus, wherein the plant virus is a comovirus.

CLPR:

4. Virus particles according to claim 1, in which the foreign peptide is inserted in the .beta.B-.beta.C loop of the plant virus.

CLPR:

6. Virus particles according to claim 1 or 5, in which the foreign peptide is incorporated in an exposed surface of the coat protein of the plant virus.

CLPR:

22. A method of producing plant virus particles according to any of claims 1 or 5, which comprises inserting a nucleotide sequence coding for a foreign peptide into the virus genome of the plant viral nucleic acid which codes for the coat protein so as to modify the plant viral nucleic acid in such a way as to avoid the production of direct sequence repeats flanking the introduced sequence, infecting plants, plant tissue, plant cells, or protoplasts with the modified viral nucleic acid, and harvesting assembled particles of said plant virus.

CLPR:

24. A method according to any of claims 22, in which the foreign nucleotide sequence is inserted by selecting two different restriction enzyme sites in the plant viral nucleic acid; cutting the plant viral nucleic acid using the corresponding restriction enzymes; and inserting into the cut viral nucleic acid a pair of complementary oligonucleotides which encode the foreign peptide and which terminate in ends compatible with the restriction enzyme cutting sites, and wherein in the complementary oligonucleotides, the sequence encoding the foreign peptide is flanked by plant virus-specific sequences so that the foreign nucleotide sequence is inserted as an addition to the plant viral nucleic acid.

CLPR:

25. A method according to claim 22, applied to an RNA plant virus, which comprises introducing a DNA sequence coding for the foreign peptide into a cDNA corresponding to the RNA of the plant virus which codes for an exposed portion of its coat protein, producing from the thus modified cDNA an RNA transcript thereof, inoculating plants, plant tissue, plant cells, or protoplasts with the transcript, optionally together with any other RNA required for

multiplication and
assembly of whole virus particles in the plant material, and harvesting
assembled particles of
the modified virus.

CLPR:

26. A method according to claim 25, in which the modified cDNA is
produced by introducing the DNA
encoding the foreign peptide into a DNA fragment excised from the plant
viral cDNA, and
recombining the modified fragment so as to reconstitute the plant viral
cDNA in modified form.

14. Document ID: US 5959181 A

L9: Entry 14 of 52

File: USPT

Sep 28, 1999

US-PAT-NO: 5959181

DOCUMENT-IDENTIFIER: US 5959181 A

TITLE: Method of preparation of transgenic plants resistant to viral
infections and so obtained
plants

DATE-ISSUED: September 28, 1999

US-CL-CURRENT: 800/301; 435/320.1, 435/410, 435/419, 435/468,
536/23.72, 536/24.1, 536/24.5,
800/278, 800/279, 800/280, 800/286, 800/295, 800/298

APPL-NO: 8/ 854170

DATE FILED: May 9, 1997

IN: Cellini; Francesco, Grieco; Pasquale Domenico

AB: The present invention relates to a method of preparation of
transgenic plants
resistant to viral infections by introducing into the genome of a host plant
an antisense
gene construct constituted by: the domain F of the subgenomic promoter
of a viral RNA; a
leader sequence of a viral ORF, downstream from said subgenomic
promoter; the gene encoding
a viral coat protein, downstream from said leader sequence; and the
3'-terminal region of a
viral RNA, downstream from said gene. The present invention also relates
to a recombinant
vector comprising a promoter functional in a host plant, and, operably
linked to this
promoter, the antisense gene construct of the present invention.

L9: Entry 14 of 52

File: USPT

Sep 28, 1999

DOCUMENT-IDENTIFIER: US 5959181 A

TITLE: Method of preparation of transgenic plants resistant to viral
infections and so obtained
plants

BSPR:

Recently, it was demonstrated for different viral species with tripartite or
monopartite RNA,
that, through the introduction into plants of RNA-4 gene encoding the coat
protein, transgenic
plants can be obtained which display a strong decrease in disease
symptoms when they are exposed

to infections with the same virus.

BSPR:

Another strategy used in order to obtain transgenic plants resistant to
determined viruses
consists in inserting into the plant a DNA sequence which is
complementary to a portion of the
viral genome in antisense orientation (not encoding). Unfortunately, the use
of such a strategy
gave unsatisfactory results. In fact, transgenic tobacco plants, obtained by
using the antisense
gene of the coat protein of CMV and PVX, displayed a tolerance to virus
only when they were
infected with low inoculum concentrations (Cuozzo et al., Biotechnol.,
6:549-557, (1988);
Hemenway et al., EMBO J. 7:1273-1280, 1988). Furthermore,
discouraging results were obtained when
antisense genes were used which were capable of complementing with
different domains of genomic
RNAs of CMV. In fact, only in one case low resistance levels were
observed (Rezaian et al., Plant
Molecular Biology, 11:463-471, 1988).

BSPR:

It has now been found that the drawbacks displayed by the prior art, as
discussed hereinabove,
can be overcome by means of the method according to the present
invention, which is based on the
use of an antisense gene construct which allows transgenic plants to be
obtained which display a
complete resistance to virus, in absence of production of coat protein.

15. Document ID: US 5955647 A

L9: Entry 15 of 52

File: USPT

Sep 21, 1999

US-PAT-NO: 5955647

DOCUMENT-IDENTIFIER: US 5955647 A

TITLE: Method for using tobacco mosaic virus to overproduce peptides
and proteins

DATE-ISSUED: September 21, 1999

US-CL-CURRENT: 800/298; 435/235.1, 435/236, 435/69.3, 530/412,
536/23.72, 800/288, 800/317.3

APPL-NO: 8/ 687559

DATE FILED: November 18, 1996

PARENT-CASE:

This application is a Continuation-in-Part application of U.S. Ser. No.
08/192,477, filed Feb. 3,
1994, now abandoned.

PCT-DATA:

APPL-NO

DATE-FILED

PUB-NO

PUB-DATE

371-DATE

102(E)-DATE

PCT/US95/01467

February 3, 1995

WO95/21248

Aug 10, 1995

Nov 18, 1996

Nov 18, 1996

IN: Fitchen; John H., Beachy; Roger N.

AB: The invention describes an infectious modified Tobacco Mosaic Virus (TMV) virion comprising a modified coat protein (CP) having a heterologous peptide inserted between amino acid residues 154 and 155 of CP. Also described is an infectious TMV virion having a modified movement protein (MP). The invention further describes nucleotide sequences encoding the modified TMV virion with either a modified CP or modified MP, and methods for producing the heterologous peptide in plants using the nucleotide sequences and modified virions.

L9: Entry 15 of 52

File: USPT

Sep 21, 1999

DOCUMENT-IDENTIFIER: US 5955647 A
TITLE: Method for using tobacco mosaic virus to overproduce peptides and proteins

BSPR:
A method is provided for overexpression in plants of heterologous peptides of from about 5 to 20 amino acids as fusion proteins inserted near, but preferably not at the carboxy terminus of the coat protein of tobacco mosaic virus using a modified cDNA infectious clone of TMV. Despite insertion of the foreign peptide sequence into the viral coat protein, stable virions are provided by this invention so that systemic infections is readily achieved in suitable plants. In some instances, the method utilizes coinfection of the plant with 1) the infectious clone having a modified CP gene, but a wild type movement protein (MP) gene and 2) a second TMV infectious clone having a wild type coat protein gene and a MP gene that has been modified to render the movement protein dysfunctional.

16. Document ID: US 5939541 A

L9: Entry 16 of 52

File: USPT

Aug 17, 1999

US-PAT-NO: 5939541
DOCUMENT-IDENTIFIER: US 5939541 A
TITLE: Method for enhancing expression of a foreign or endogenous gene product in plants
DATE-ISSUED: August 17, 1999

US-CL-CURRENT: 536/24.1; 435/320.1, 435/411, 435/468, 536/23.72, 800/287, 800/288

APPL-NO: 8/ 827575
DATE FILED: March 28, 1997

IN: Vance; Vicki B., Pruss; Gail J., Dawson; William O., Carrington; James, Marton; Laszlo

AB: The present invention provides a method for enhancing the expression of genes in plants by supplying a virally encoded booster sequence comprising the 5'

proximal region of the poliovirus genome to the plant. The booster sequence enhances the expression of foreign genes or endogenous plant genes in plants by employing any known methodology of expressing introduced genes in plants. The booster sequence may be used to enhance expression of any gene, including foreign genes or endogenous plant genes, introduced by means of stable transformation into the genome of the plant or introduced by expression from a plant viral expression vector.

L9: Entry 16 of 52

File: USPT

Aug 17, 1999

DOCUMENT-IDENTIFIER: US 5939541 A
TITLE: Method for enhancing expression of a foreign or endogenous gene product in plants

DEPR:
The present Example sets forth an exemplary method for enhancing expression of an endogenous plant gene or a foreign gene (or a portion of a foreign or endogenous gene) that has been introduced to a plant as a fusion to a viral protein expressed from a viral vector. A viral vector expressing a foreign gene or an endogenous plant sequence as a fusion to the coat protein of the virus, such as the vector described in Sugiyama, Hamamoto, Takemoto, Watanabe, Okada, "Systematic Production of Foreign Peptides on the Particle Surface of Tobacco Mosaic Virus," 359 FEBS Lett., 247 250 (1995), is one such example. The viral vector may be used to infect a transgenic plant host that supplies the booster sequence via expression from stably incorporated DNA copies of said booster sequence, for example the U-6B transgenic tobacco plants described herein. The expression of the foreign peptides fused to the viral coat protein would be enhanced.

17. Document ID: US 5919457 A

L9: Entry 17 of 52

File: USPT

Jul 6, 1999

US-PAT-NO: 5919457
DOCUMENT-IDENTIFIER: US 5919457 A
TITLE: TXU-5/B53-PAP antiviral biotherapeutic agent for the treatment of AIDS
DATE-ISSUED: July 6, 1999

US-CL-CURRENT: 424/183.1; 424/160.1, 435/339.1, 435/70.21, 530/370, 530/379, 530/388.35, 530/389.4, 530/391.9

APPL-NO: 8/ 584966
DATE FILED: January 11, 1996

IN: Uckun; Fatih M.

AB: Immunoconjugates comprising the monoclonal antibody TXU-5/B53 linked to pokeweed antiviral protein or bioactive subunits thereof are provided which are

effective for the
treatment of mammalian HIV infection.

L9: Entry 17 of 52

File: USPT

Jul 6, 1999

DOCUMENT-IDENTIFIER: US 5919457 A
TITLE: TXU-5/B53-PAP antiviral biotherapeutic agent for the treatment of
AIDS

DEPR:

Pokeweed antiviral protein (PAP) is an antiviral agent isolated from the
leaves or seeds of
Phytolacca americana (Irvin and Uckun, Pharmacology and Therapeutics
55: 279, 1992). PAP displays
broad-spectrum antiviral activity against plant viruses, herpes simplex
virus, cytomegalovirus,
poliovirus, and influenza virus. Aron et al., Agents Chemotherapy, 17,
1032 (1980). In fact,
pokeweed antiviral protein was discovered due to its ability to inhibit the
transmission of
tobacco mosaic virus (TMV) in plants and it was subsequently
demonstrated that the purified
protein was equally effective against a number of other plant viruses.
Tomlinson et al., J. Gen.
Virol., 22, 225 (1974).

18. Document ID: US 5907084 A

L9: Entry 18 of 52

File: USPT

May 25, 1999

US-PAT-NO: 5907084
DOCUMENT-IDENTIFIER: US 5907084 A
TITLE: Virus resistant or tolerant cells
DATE-ISSUED: May 25, 1999

US-CL-CURRENT: 800/279; 435/320.1, 435/411, 435/414, 536/23.1,
536/24.5, 800/280, 800/301

APPL-NO: 8/ 624581
DATE FILED: April 3, 1996

FOREIGN-APPL-PRIORITY-DATA:
COUNTRY

	APPL-NO	APPL-DATE
GB	9320548	October 6, 1993

PCT-DATA:
APPL-NO
DATE-FILED
PUB-NO
PUB-DATE
371-DATE
102(E)-DATE
PCT/EP94/03295
October 5, 1994
WO95/09920
Apr 13, 1995
Apr 3, 1996
Apr 3, 1996

IN: de Haan; Petrus Theodorus

AB: A nucleotide sequence comprising a transcriptional regulatory
sequence and a
sequence contiguous therewith and under the transcriptional control
thereof, which
contiguous sequence encodes an RNA which consists of a plurality of
sub-sequences,
characterized in that at least two of the sub-sequences have the sequences
of viral RNAs and
the RNA contains at least one translational stop codon located upstream
of the 3' terminal
sub-sequence. It is preferred that at least one of the sub-sequences is in an
anti-sense
configuration with respect to virus RNA, and that the contiguous
sequence encodes mRNA. The
invention also includes, inter alia, the use of such a sequence in the
generation of virus
resistant or tolerant plants, and such plants comprising the sequence.

L9: Entry 18 of 52

File: USPT

May 25, 1999

DOCUMENT-IDENTIFIER: US 5907084 A
TITLE: Virus resistant or tolerant cells

BSPR:

Numerous attempts have been made to engineer viral resistance into plants
by inserting
DNA-containing vectors into acceptor plant tissue, which DNA is capable
of encoding viral
proteins in the thus transformed plant. The viral protein may confer
resistance to an invading
virus comprising a viral protein substantially the same as that encoded by
the introduced DNA.
Other attempts at engineering virus resistance in plants use anti-sense RNA
which relies on the
introduction of DNA encoding an RNA strand which is complementary to
the RNA of an invading virus
and thus interferes with the replication thereof. Plants displaying a broad
degree of reduced
susceptibility, i.e. to more than one viral type, or a greater degree of
reduced susceptibility
to a particular virus type, are highly desirable.

DEPR:

(1). Multigene DNA constructs comprising at least one non-structural gene
wherein the multigene
DNA is under the control of a single set of genetic regulatory elements. By
"non-structural gene"
is meant a gene capable of coding for a viral RNA molecule which is
substantially incapable of
encoding for a viral polypeptide or protein but which is nevertheless
capable of conferring an
RNA mediated reduced susceptibility of a plant virus in plant cells. (2).
Constructs according to
clause 1, wherein the at least one non-structural gene is a viral gene. (3).
Multigene DNA
constructs comprising at least one non-structural gene wherein the
multigene DNA constructs are
capable of giving rise to viral elements in plant cells which are capable of
conferring a reduced
susceptibility to plant viruses in plant cells, and wherein the multigene
DNA is under the
control of a single set of genetic regulatory elements. (4) Constructs
according to clauses 2 to
3 comprising at least one viral non-structural gene and one viral structural
gene. (5) Constructs
according to any one of clauses 2 to 4 comprising at least two
non-structural genes and no viral
structural gene elements. (6). Constructs according to clause 5 comprising
from 2 to 5 viral
non-structural genes. (7). Constructs according to any one of clauses 1 to 3

comprising DNA

capable of coding for viral RNA molecules of tospoviruses, potyviruses, potexviruses,

tobamoviruses, luteoviruses, cucumoviruses, bromoviruses, closteroviruses, tombusviruses, and

furoviruses. (8) Constructs according to clause 7, wherein the DNA codes for non-structural viral

RNA molecules of nucleocapsid proteins, viral coat proteins, and non-structural viral proteins.

(9). Plants comprising multigene DNA constructs of any one of clauses 1 to 3. (10). Plants

according to clause 9 selected from the group comprising tomatoes, peppers, melons, lettuces,

cauliflowers, broccolis, cabbages, brussels sprouts, sugar beet, corn (maize), sweetcorn, onions,

carrots, leeks, cucumbers, tobacco's alfalfa's, aubergines, beets, broad beans, celery's,

chicory's, cow peas, endives, gourds, groundnuts, papayas, peas, peanuts, pineapples, potatoes,

safflowers, snap beans, soybeans, spinaches, squashes, sunflowers, water-melons, and sorghums.

(11). Plants according to clause 9 selected from the group ornamentals consisting essentially of

Impatiens, IBegonias, Petumias, Pelargoniums (geraniums), Violas, Cyclamens, Verbenas, Vincas,

Tagetes, Primulas, Saint Paulia's Ageratums, Amaranthuses, Anthirrhinums, Aquilegias,

Chrysanthemums, Cineraria, Clovers, Cosmos's, Cowpeas, Dahlia's, Daturas, Delphiniums, Gerbera's,

Gladioluses, Gloxinias, Hippeastrums, Mesembryanthemums, Salpiglossis, and Zinnias. (12). A

method for obtaining plants displaying a reduced susceptibility to viruses which comprises: (a)

inserting into the genome of a plant cell a DNA construct according to any one of clauses 1 to 8;

(b) obtaining transformed cells; and (c) regenerating from the transformed cells genetically

transformed plants.

19. Document ID: US 5874087 A

L9: Entry 19 of 52

File: USPT

Feb 23, 1999

US-PAT-NO: 5874087

DOCUMENT-IDENTIFIER: US 5874087 A

TITLE: Modified plant viruses as vectors

DATE-ISSUED: February 23, 1999

US-CL-CURRENT: 424/199.1; 424/185.1, 424/186.1, 424/192.1, 424/202.1, 424/204.1, 435/235.1

APPL-NO: 8/ 471048

DATE FILED: June 6, 1995

PARENT-CASE:

This application is a Division of application Ser. No. 08/137,032, filed as PCT/GB92/00589, Apr. 2, 1992.

FOREIGN-APPL-PRIORITY-DATA:

COUNTRY

APPL-NO

APPL-DATE

GB

91 08386

April 19, 1991

IN: Lomonosoff; George Peter, Johnson; John Emil

AB: The invention relates to assembled particles of a plant virus containing a

predetermined foreign peptide as part of the coat protein of the virus, and a method for

their production. The foreign peptide is preferably a biologically functional peptide, the

biological application of which requires or is enhanced by presentation of the peptide in

association with a larger molecule or particle.

L9: Entry 19 of 52

File: USPT

Feb 23, 1999

DOCUMENT-IDENTIFIER: US 5874087 A

TITLE: Modified plant viruses as vectors

ABPL:

The invention relates to assembled particles of a plant virus containing a predetermined foreign

peptide as part of the coat protein of the virus, and a method for their production. The foreign

peptide is preferably a biologically functional peptide, the biological application of which

requires or is enhanced by presentation of the peptide in association with a larger molecule or

particle.

BSPR:

This invention relates to the use of viruses as carriers (vectors) for the production or

presentation of foreign peptides. More particularly, the invention relates to the genetic

manipulation of viral nucleic acid by incorporation of foreign nucleic acid sequences which are

expressed as peptides in the virus particle (virion). In this specification the term "foreign",

as applied to a peptide or to the nucleic acid encoding therefor, signifies peptide or nucleic

acid sequences which are not native to the plant virus used as a vector.

Such sequences can be

alternatively described as exogenous or heterologous sequences. The term "peptide" includes small

peptides and polypeptides.

BSPR:

The present invention utilises plant viruses as vector systems for the expression of foreign

nucleotide sequences ie nucleotide sequences (RNA or DNA) which are not present in plant viruses,

as found in Nature, and which in consequence code for peptides not normally found in any

naturally occurring plant virus.

BSPR:

The present invention comprises assembled particles of a plant virus containing a foreign

peptide. The plant viruses of the present invention are therefore modified forms of the native

viruses and for convenience will be referred to as modified viruses.

BSPR:

The foreign peptides which may be incorporated into plant viruses according to this invention may

be of highly diverse types and are subject only to the limitation that the nature and size of the

foreign peptide and the site at which it is placed in or on the virus particle do not interfere

with the capacity of the modified virus to assemble when cultured in vitro or in vivo. In broad

concept, modified viruses may be formed from any biologically useful peptides (usually

polypeptides) the function of which requires a particular conformation for its activity. This may

be achieved by association of the peptide with a larger molecule eg to improve its stability or mode of presentation in a particular biological system. Examples of such peptides are peptide hormones; enzymes; growth factors; antigens of protozoal, viral, bacterial, or fungal origin; antibodies including anti-idiotypic antibodies; immunoregulators and cytokines eg interferons and interleukins; receptors; adhesins; and parts or precursors of any of the foregoing types of peptide. The peptide preferably contains more than 5 amino acids.

BSPR:
To produce modified plant virus particles in accordance with this invention the plant viral nucleic acid is modified by introducing a nucleotide sequence coding for the foreign peptide eg an animal virus antigen at that part of the plant viral genome which codes for an exposed portion of the coat protein, infecting plants or plant cells with the modified viral nucleic acid, and harvesting assembled particles of the modified virus. This procedure is best carried out by direct manipulation of the DNA of the virus in the case of DNA viruses or by manipulation of a cDNA corresponding to the RNA of an RNA virus. In the case of an RNA virus, an RNA transcript of the modified DNA is usually prepared for inoculation of plant cells or preferably whole plants so as to achieve a multiplication stage prior to the harvesting of assembled particles of the modified virus. In the case of a DNA virus, the DNA itself is introduced into the plant. In this way, the foreign peptide is initially expressed as part of the capsid protein and is thereby produced as part of the whole virus particle. The peptide may thus be produced as a conjugate molecule intended for use as such. Alternatively, the genetic modification of the virus may be designed in order to permit release of the desired peptide by the application of appropriate agents which will effect cleavage from the virus particle.

CLPR:
3. A method according to claim 1, which comprises (a) introducing a DNA sequence coding for the foreign peptide into a cDNA corresponding to the plant virus RNA which codes for an exposed portion of the plant virus coat protein, (b) producing from the thus modified cDNA an RNA transcript thereof, (c) inoculating a member of the group consisting of plants, plant tissue, plant cells, and protoplasts with the transcript, if necessary together with any other RNA required for multiplication and assembly of whole virus particles in the plant material, and (d) harvesting assembled particles of the virus.

CLPR:
6. A method according to claim 1, which comprises (a) introducing a DNA sequence coding for the foreign peptide into a cDNA corresponding to the plant virus RNA which codes for an exposed portion of the plant virus coat protein; (b) inoculating a member of the group consisting of plants, plant tissue, plant cells, and protoplasts with the thus modified cDNA, if necessary together with any other RNA required for multiplication and assembly of whole virus particles in the plant material; and (c) harvesting assembled particles of the virus.

20. Document ID: US 5830887 A

L9: Entry 20 of 52

File: USPT

Nov 3, 1998

US-PAT-NO: 5830887
DOCUMENT-IDENTIFIER: US 5830887 A
TITLE: Health supplements containing phyto-oestrogens, analogues or metabolites thereof
DATE-ISSUED: November 3, 1998

US-CL-CURRENT: 514/182; 424/423, 424/449, 424/451, 424/464, 424/757, 426/545, 514/25, 549/403, 549/406

APPL-NO: 8/ 338567
DATE FILED: January 12, 1995

FOREIGN-APPL-PRIORITY-DATA:
COUNTRY

	APPL-NO	APPL-DATE
AU	PL2511	May 19, 1992

PCT-DATA:			
APPL-NO	DATE-FILED	PUB-NO	PUB-DATE
			371-DATE
			102(E)-DATE
PCT/AU93/00230			
May 19, 1993			
	WO93/23069		
	Nov 25, 1993		
	Jan 12, 1995		
	Jan 12, 1995		

IN: Kelly; Graham Edmund

AB: Compositions enriched with natural phyto-oestrogens or analogues thereof selected from Genistein, Daidzein, Formononetin and Biochanin A. These may be used as food additives, tablets or capsules for promoting health in cases of cancer, pre-menstrual syndrome, menopause or hypercholesterolaemia.

L9: Entry 20 of 52

File: USPT

Nov 3, 1998

DOCUMENT-IDENTIFIER: US 5830887 A
TITLE: Health supplements containing phyto-oestrogens, analogues or metabolites thereof

BSPR:
There are 3 principal classes of phyto-oestrogens, viz. isoflavones, lignans, and coumestans. The isoflavones are thought to have a broad range of biological functions in plants, although these are poorly understood. However, two particular functions are recognised--(a) as phyto-alexin or stressor chemicals which are secreted by the plant in response to attack by parasites such as insects, fungi, viruses, etc. and which display activity against these parasites, and (b) chemicals which encourage colonisation of nitrogen-fixing bacteria on the roots of legumes. The

biological functions in plants of the lignans and coumestans is not generally understood.

21. Document ID: US 5824857 A

L9: Entry 21 of 52

File: USPT

Oct 20, 1998

US-PAT-NO: 5824857
DOCUMENT-IDENTIFIER: US 5824857 A
TITLE: Plant promoter
DATE-ISSUED: October 20, 1998

US-CL-CURRENT: 800/287; 435/410, 435/419, 536/23.4, 536/23.6, 536/24.1, 800/293, 800/320, 800/320.2

APPL-NO: 7/ 789738
DATE FILED: November 8, 1991

IN: Beachy; Roger N., Bhattacharyya; Maitrayee

AB: A genome length transcript promoter from a badnavirus, rice tungro bacilliform virus (RTBV), is disclosed and its DNA sequence provided. This promoter drives expression specifically in vascular tissues of plants. This promoter sequence may be utilized in a chimeric gene to drive the tissue specific expression of a foreign structural gene in vascular tissue of transgenic plants.

L9: Entry 21 of 52

File: USPT

Oct 20, 1998

DOCUMENT-IDENTIFIER: US 5824857 A
TITLE: Plant promoter

BSPR:
Even though providing constitutive expression of a gene in plants is often desirable, it is also desirable in some instances to direct expression of a gene to particular tissues in a plant. Some tissue specific plant promoters are known, such as those capable of directing expression preferentially in the fruit of a plant, but the genome length transcript, i.e. full length transcript or major transcript, promoters that have been obtained from double-stranded DNA plant viruses all display strong, constitutive expression patterns. Extensive structure analysis of the CaMV35S promoter sequence has identified domains within the viral promoter sequence that confer tissue specificity (Odell et al. 1985; Benfey et al. 1989; Fang et al. 1989), but the promoter sequence in its entirety is a constitutive promoter.

22. Document ID: US 5633434 A

L9: Entry 22 of 52

File: USPT

May 27, 1997

US-PAT-NO: 5633434
DOCUMENT-IDENTIFIER: US 5633434 A
TITLE: Transgenic plants displaying virus and phosphinothricin resistance
DATE-ISSUED: May 27, 1997

US-CL-CURRENT: 800/280; 435/193, 435/418, 435/419, 435/69.1, 504/207, 536/23.1, 536/23.2, 536/23.72, 800/300, 800/301

APPL-NO: 8/ 279706
DATE FILED: July 25, 1994

PARENT-CASE:
This application is a continuation-in-part, continuation of application Ser. No. 08/123,699, filed Sep. 17, 1993, now abandoned, which in turn is a continuation of application Ser. No. 07/910,329, filed as PCT/EP91/00130, Jan. 24, 1991, which in turn is abandoned.

FOREIGN-APPL-PRIORITY-DATA:
COUNTRY

APPL-NO	APPL-DATE
DE 4003045	February 2, 1990

IN: Schneider; Rudolf, Donn; G unter, M ullner; Hubert

AB: Virus genes, for example coat protein genes, which bring about a reduction in the signs of infection by the corresponding virus or bring about virus resistance can be combined with herbicide-resistance genes for the transformation of plants. A combination of this type facilitates the selection of the transgenic plants. In addition, in practical field cultivation, the vitality of the plants is increased by the virus tolerance, and an improved plant protection is possible owing to the herbicide-resistance gene.

L9: Entry 22 of 52

File: USPT

May 27, 1997

DOCUMENT-IDENTIFIER: US 5633434 A
TITLE: Transgenic plants displaying virus and phosphinothricin resistance

23. Document ID: US 5618699 A.

L9: Entry 23 of 52

File: USPT

Apr 8, 1997

US-PAT-NO: 5618699
DOCUMENT-IDENTIFIER: US 5618699 A
TITLE: Plant virus vector, plasmid, process for expression of foreign gene and process for obtaining foreign gene product
DATE-ISSUED: April 8, 1997

US-CL-CURRENT: 435/69.7; 435/235.1, 435/320.1, 435/69.1, 435/70.1, 536/23.72

APPL-NO: 8/ 313127

DATE FILED: November 30, 1994

FOREIGN-APPL-PRIORITY-DATA:
COUNTRY

	APPL-NO	APPL-DATE
JP	4-108628	March 31, 1992
JP	4-188744	June 22, 1992
JP	4-351970	December 8, 1992

PCT-DATA:
APPL-NO

	DATE-FILED	PUB-NO	PUB-DATE	371-DATE	102(E)-DATE
PCT/JP93/00408	March 31, 1993	WO93/20217	Oct 14, 1993	Nov 30, 1994	Nov 30, 1994

IN: Hamamoto; Hiroshi, Sugiyama; Yoshinori, Nakagawa; Noriaki, Hashida; Eiji, Tsuchimoto; Suguru, Nakanishi; Noriyuki, Matsunaga; Yuji, Okada; Yoshimi

AB: The present invention relates to a plant virus vector comprising a foreign gene linked downstream of a coat protein gene of tobacco mosaic virus via a nucleotide sequence which cause the readthrough, and a plasmid which is transcribed to provide the vector, as well as a process for expression of a foreign gene in a plant by inoculating the plant with the vector. In addition, the present invention relates to a process for efficiently recovering a foreign gene product produced in a plant as virions.

L9: Entry 23 of 52

File: USPT

Apr 8, 1997

DOCUMENT-IDENTIFIER: US 5618699 A

TITLE: Plant virus vector, plasmid, process for expression of foreign gene and process for obtaining foreign gene product

BSPR:

Since the present plant virus vector has a nucleotide sequence which causes readthrough, it simultaneously produces both a wild type coat protein and a fused protein (i.e., a fused protein comprising a desired protein or peptide derived from a foreign gene and the coat protein).

Therefore, virions normally result in systemic infection and expression of the foreign gene throughout a whole plant.

24. Document ID: US 5589625 A

L9: Entry 24 of 52

File: USPT

Dec 31, 1996

US-PAT-NO: 5589625
DOCUMENT-IDENTIFIER: US 5589625 A
TITLE: Transgenic plants displaying multiple virus resistance and a process for their production
DATE-ISSUED: December 31, 1996

US-CL-CURRENT: 800/279; 435/418, 435/69.1, 800/301

APPL-NO: 8/ 374229

DATE FILED: January 18, 1995

PARENT-CASE:

This application is a Continuation-in-Part application of U.S. Ser. No. 07/965,343, filed Oct. 23, 1992, now abandoned.

FOREIGN-APPL-PRIORITY-DATA:
COUNTRY

	APPL-NO	APPL-DATE
EP	92104676	March 18, 1992

IN: Saarma; Mart, Kelve; Merikke, Truve; Erkki, Teeri; Teemu

AB: This invention discloses transgenic plants, such as transgenic tobacco and potato, having resistance to multiple viral taxonomic groups using parts of the 2,5A oligoadenylate pathway. In particular, said plants are genetically engineered to contain a DNA sequence encoding at least one polypeptide having a 2,5A synthetase activity. By this means a step in the 2,5A oligoadenylate pathway heretofore believed to be missing in all plants is provided so that viral infection in the transgenic plants is inhibited via a 2,5A dependent endonuclease. Moreover, this invention relates to a process for the production of said transgenic plants by transfection with a genetically engineered DNA sequence encoding at least one polypeptide having a 2,5A synthetase activity.

L9: Entry 24 of 52

File: USPT

Dec 31, 1996

DOCUMENT-IDENTIFIER: US 5589625 A
TITLE: Transgenic plants displaying multiple virus resistance and a process for their production

BSPR:

In summary, the prior art does not permit the conclusion that the 2,5A oligoadenylate pathway can be used as a basis for constructing transgenic plants displaying multiple virus resistance. Thus, the technical problem of the present invention is to provide a transgenic plant displaying resistance to multiple virus taxonomic groups using parts of the 2,5A oligoadenylate pathway.

BSPR:

One object of the present invention relates to a transgenic plant displaying multiple virus resistance which contains a genetically engineered DNA sequence encoding at least one polypeptide having a 2,5A synthetase activity, wherein said polypeptide upon expression is capable of activating an endonuclease causing degradation of viral RNA.

BSPR:

In the present invention a transgenic plant is provided that displays resistance to multiple viral taxonomic groups. The transgenic plant contains a genetically engineered DNA sequence encoding at least one polypeptide having a 2,5A synthetase activity, wherein the polypeptide activates an endonuclease contained within the plant, thereby causing degradation of viral RNA so as to prevent or lessen infection. The invention also discloses propagating material derived from such transgenic plants.

DEPR:

This invention provides a method for obtaining transgenic plants displaying resistance to multiple viral taxonomic groups by restoring to said plants a functioning 2,5A oligoadenylate pathway. In particular, plants containing parts of the 2,5A oligoadenylate pathway are genetically engineered to contain a DNA sequence encoding at least one polypeptide having a 2,5A synthetase activity. By this means a step in the 2,5A oligoadenylate pathway heretofore believed to be missing in all plants is provided, so that viral infection in the transgenic plants is inhibited via a 2,5A dependent endonuclease. Moreover, this invention relates to a process for the production of said transgenic plants by transfection with a genetically engineered DNA sequence encoding at least one polypeptide having a 2,5A synthetase activity.

DEPR:

A further object of the present invention is a propagating material derived from a transgenic plant displaying resistance to multiple viral taxonomic groups.

DEPR:

A further object of the present invention is a process for the production of a transgenic plant displaying resistance to multiple viral taxonomic virus groups comprising the introduction of a genetically engineered DNA sequence encoding at least one polypeptide having a 2,5A synthetase activity, into the genetic material of a suitable plant.

DEPR:

A further object of the present invention is the use of a genetically engineered DNA sequence encoding at least one polypeptide having a 2,5A synthetase activity, for the production of a transgenic plant displaying resistance to multiple viral taxonomic groups.

CLPR:

1. A transgenic plant that utilizes the 2,5A oligoadenylate pathway and displays resistance to multiple plant viral taxonomic groups comprising a genetically engineered DNA sequence encoding at least one polypeptide having a 2,5A synthetase activity, wherein said polypeptide activates an endonuclease causing degradation of viral RNA.

CLPR:

17. A transgenic plant cell that utilizes the 2,5A oligoadenylate pathway and displays resistance to multiple plant virus taxonomic groups comprising a genetically engineered DNA sequence encoding at least one polypeptide having a 2,5A synthetase activity.

CLPR:

23. A plant cell displaying resistance to multiple plant virus taxonomic groups descended from the plant cell of claim 17 and comprising said 2,5A synthetase activity.

ORPL:

Truve, et al., Principles and background for the construction of transgenic plants displaying multiple virus resistance, Arch Virol (1994) [Suppl] 9:41-50.

25. Document ID: US 5583021 A

L9: Entry 25 of 52

File: USPT

Dec 10, 1996

US-PAT-NO: 5583021

DOCUMENT-IDENTIFIER: US 5583021 A

TITLE: Production of virus resistant plants

DATE-ISSUED: December 10, 1996

US-CL-CURRENT: 800/280; 435/252.3, 435/320.1, 435/418, 435/419, 435/468, 435/469, 435/470, 536/23.72, 800/301

APPL-NO: 8/ 271829

DATE FILED: July 7, 1994

PARENT-CASE:

CROSS REFERENCE TO RELATED APPLICATION This application is a continuation-in-part of application Ser. No. 07/838,509, filed Feb. 19, 1992, now abandoned.

IN: Dougherty; William G., Lindbo; John A.

AB: A method of suppressing virus gene expression in plants using untranslatable plus sense RNA is disclosed. The method is useful for the production of plants that are resistant to virus infection.

L9: Entry 25 of 52

File: USPT

Dec 10, 1996

DOCUMENT-IDENTIFIER: US 5583021 A

TITLE: Production of virus resistant plants

DEPR:

All transgenic plant lines tested displayed wild-type sensitivities to PVY and to cucumber mosaic virus. Typical necrotic local lesions formed when the 2RC lines were inoculated with tobacco mosaic virus (data not shown).

DEPR:

While not wishing to be bound by speculation, it is suggested that the highly resistant and recovery phenotypes of the 2RC lines can be accommodated by the following working model. This model suggests the existence of an inducible, cytoplasmic-based, cellular activity that degrades specific RNA sequences. In transgenic plants displaying the recovery phenotype, this RNA degradation system is activated only after virus infection and by the additive level of transgene RNA and viral RNA present. In contrast, the highly resistant lines may

have the activity fully induced by the transgene transcript. The failure of a rootstock from a highly resistant line to induce a scion from a susceptible line in grafting studies suggests the activity is a programmed cell response not induced via a diffusible signaling molecule as is the case with systemically acquired resistance (Kuc 1982; Ward et al., 1991). Once the antiviral system is activated, it is absolute in its efficacy against TEV, yet it is not effective against the closely related virus PVY.

DEPR:

The engineered resistance was TEV specific. None of the lines tested displayed any resistance to PVY or tomato spotted wilt virus (TSWV) as a limited number of plants were naturally infected with these viruses in the plot over the 2 year study.

DEPV:

Susceptible plant: A plant that supports viral replication and displays virus-induced symptoms.

26. Document ID: US 5510253 A

L9: Entry 26 of 52

File: USPT

Apr 23, 1996

US-PAT-NO: 5510253
DOCUMENT-IDENTIFIER: US 5510253 A
TITLE: Plants resistant to infection by PLRV
DATE-ISSUED: April 23, 1996

US-CL-CURRENT: 800/279; 435/320.1, 435/69.1, 536/23.72, 800/301

APPL-NO: 8/ 326297
DATE FILED: October 20, 1994

PARENT-CASE:

This is a Continuation of application Ser. No. 08/012,688, filed Feb. 3, 1993 now abandoned.

IN: Mitsky; Timothy A., Hemenway; Cynthia L., Turner; Nilgun E.

AB: An isolated DNA sequence which codes for a PLRV replicase gene is disclosed herein. A method for providing resistance to infection by a virus by expressing a replicase gene in plants is also disclosed, as are transgenic potato plants and tubers containing the replicase gene.

L9: Entry 26 of 52

File: USPT

Apr 23, 1996

DOCUMENT-IDENTIFIER: US 5510253 A
TITLE: Plants resistant to infection by PLRV

DEPR:

PLRV is not mechanically transmissible. Spread of PLRV from infected plants to uninfected plants can only be accomplished by aphids. This is the reason that insecticide application is currently necessary for controlling this disease. Virus resistant potatoes will no

longer require insecticides to control aphids. Potato cultivars which are resistant to PLRV often display no or reduced titers of PLRV and the virus is not as easily transmitted from these plants to other plants by aphids. This characteristic is of commercial significance because it limits the potential for virus epidemics in the field.

27. Document ID: US 5376675 A

L9: Entry 27 of 52

File: USPT

Dec 27, 1994

US-PAT-NO: 5376675
DOCUMENT-IDENTIFIER: US 5376675 A
TITLE: Control of parasitic nematodes (A)
DATE-ISSUED: December 27, 1994

US-CL-CURRENT: 514/425

APPL-NO: 8/ 070391
DATE FILED: August 30, 1993

FOREIGN-APPL-PRIORITY-DATA:
COUNTRY

APPL-NO	APPL-DATE
GB 9026271	December 3, 1990

PCT-DATE: APPL-NO	DATE-FILED	PUB-NO	PUB-DATE	371-DATE	102(E)-DATE
PCT/GB91/02111	November 28, 1991	WO92/09202	Jun 11, 1992	Aug 30, 1983	Aug 30, 1993

IN: Alphey; Thomas J. W., Birch; Andrew N. E., Fellows; Linda E., Robertson; Walter M.

AB: The use of the compound 2R,5R-dihydroxymethyl-3R,4R-dihydroxypyrrolidine (DMDP) ##STR1## or an acid addition salt thereof in controlling diseases caused by parasitic nematodes in plants or mammals.

L9: Entry 27 of 52

File: USPT

Dec 27, 1994

DOCUMENT-IDENTIFIER: US 5376675 A
TITLE: Control of parasitic nematodes (A)

BSPR:

DMDP displays its properties against a wide range of nematodes affecting plants, e.g. root-knot nematodes, cyst nematodes and virus-transmitting nematodes. Of particular note is its activity against the crop-damaging nematodes of the following genera: Meloidogyne, Globodera, Heterodera, Radopholus, Pratylenchus, Hirschmanniella, Scutellonema, Helicotylenchus, Tylenchus, Rotylenchus, Ditylenchus, Longidorus, Xiphinema. With regard to nematodes which infest mammals, DMDP is active against a wide range of helminthic nematodes, especially those of the following genera: Haemonchus, Teladorsagia, Nematodirus, Trichostrongylus, Dictyocaulus and Cooperia, particularly the species Haemonchus contortus and Teladorsagia circumcincta (previously classified as Osteragia circumcincta).

28. Document ID: US 5185253 A

L9: Entry 28 of 52

File: USPT

Feb 9, 1993

US-PAT-NO: 5185253
DOCUMENT-IDENTIFIER: US 5185253 A
TITLE: Virus resistant plants
DATE-ISSUED: February 9, 1993

US-CL-CURRENT: 800/279; 435/69.1, 435/70.1, 800/294

APPL-NO: 7/ 606641
DATE FILED: October 31, 1990

PARENT-CASE:
This application is a division of U.S. Ser. No. 07/302,498 filed Jan. 27, 1989, now U.S. Pat. No. 4,970,168.

IN: Tumer; Nilgun E.

AB: Transgenic plants are disclosed which are resistant to virus infection by Potato Virus X and Potato Virus Y. Plant genes and transformation vectors are also disclosed. Potato plants, for example, Russet Burbank variety, are made resistant to dual infection by Potato Virus X and Potato Virus Y by transforming the plant to express the coat proteins of the two viruses.

L9: Entry 28 of 52

File: USPT

Feb 9, 1993

DOCUMENT-IDENTIFIER: US 5185253 A
TITLE: Virus resistant plants

DEPR:
Accordingly, the present invention provides a method for genetically engineering plants by insertion into the plant genome a DNA construct containing, inter alia, a small portion of the viral genome of PVX and PVY such that the engineered plants display resistance to the plant virus.

29. Document ID: US 4970168 A

L9: Entry 29 of 52

File: USPT

Nov 13, 1990

US-PAT-NO: 4970168
DOCUMENT-IDENTIFIER: US 4970168 A

TITLE: Virus-resistant plants

DATE-ISSUED: November 13, 1990

INVENTOR-INFORMATION:

NAME

CITY

STATE

ZIP CODE

COUNTRY

Tumer; Nilgun E.

Chesterfield

MO

N/A

N/A

US-CL-CURRENT: 800/301; 435/252.3, 435/252.33, 435/320.1, 435/417, 435/71.1

IN: Tumer; Nilgun E.

L9: Entry 29 of 52

File: USPT

Nov 13, 1990

DOCUMENT-IDENTIFIER: US 4970168 A
TITLE: Virus-resistant plants

DEPR:
Accordingly, the present invention provides a method for genetically engineering plants by insertion into the plant genome a DNA construct containing, inter alia, a small portion of the viral genome of PVX and PVY such that the engineered plants display resistance to the plant virus.

30. Document ID: JP 05328977 A

L9: Entry 30 of 52

File: JPAB

Dec 14, 1993

PUB-NO: JP405328977A
DOCUMENT-IDENTIFIER: JP 05328977 A
TITLE: PLANT VIRUS VECTOR AND PLASMID AND METHOD FOR ALLOWING TO EXPRESS FOREIGN GENE IN PLANT

PUBN-DATE: December 14, 1993

INVENTOR-INFORMATION:

NAME

COUNTRY

HAMAMOTO, HIROSHI
SUGIYAMA, YOSHINOBU
NAKANISHI, NORIYUKI
NAKAGAWA, NORIAKI

TSUCHIMOTO, TAKU
HASHIDA, HIDEJI
MATSUNAGA, YUJI
OKADA, YOSHIMI

INT-CL (IPC): C12N 15/62; C12N 15/83; C12P 21/02

IN: HAMAMOTO, HIROSHI, SUGIYAMA, YOSHINOBU,
NAKANISHI, NORIYUKI, NAKAGAWA, NORIAKI,
TSUCHIMOTO, TAKU, HASHIDA, HIDEJI, MATSUNAGA, YUJI,
OKADA, YOSHIMI

L9: Entry 30 of 52

File: JPAB

Dec 14, 1993

DOCUMENT-IDENTIFIER: JP 05328977 A
TITLE: PLANT VIRUS VECTOR AND PLASMID AND METHOD
FOR ALLOWING TO EXPRESS FOREIGN GENE IN PLANT

FPAR:

CONSTITUTION: A plant virus vector is characterized in that a foreign gene is connected to the downstream of an exodermis protein gene of a plant virus through a base sequence inducing a read-through. The exodermis protein virus is preferably the gene of tobacco virus, etc. Examples of the foreign gene includes a peptide gene having pharmacological and physiological activities, a protein gene giving stress resistance and resistance against diseases and pests to plants, and a protein gene for changing the shapes and flower colors of plants. When the virus is RNA, the plant virus vector is obtained by inserting the cDNA of a plant RNA virus into the downstream of a promoter of a plasmid having the promoter for transcription in vitro, inserting a foreign gene into the downstream of the exodermis protein gene originated from the plant virus through a base sequence inducing a read-through, and subsequently producing the RNA by a transcription reaction in vitro.

31. Document ID: WO 9911806 A1

L9: Entry 31 of 52

File: EPAB

Mar 11, 1999

PUB-NO: WO009911806A1
DOCUMENT-IDENTIFIER: WO 9911806 A1
TITLE: PATHOGEN-RESISTANT TRANSGENIC PLANTS AND
METHODS OF MAKING

PUBN-DATE: March 11, 1999

INVENTOR-INFORMATION:
NAME

COUNTRY

WANG, CHUNLIN

N/A

INT-CL (IPC): C12N 15/82; C12N 15/53; C12N 9/02; A01H 5/00
EUR-CL (EPC): C12N015/82; A01N063/02, C12N015/82, C12N015/82

IN: WANG, CHUNLIN

L9: Entry 31 of 52

File: EPAB

Mar 11, 1999

DOCUMENT-IDENTIFIER: WO 9911806 A1
TITLE: PATHOGEN-RESISTANT TRANSGENIC PLANTS AND
METHODS OF MAKING

FPAR:

The present invention provides pathogen-resistant transgenic plants and methods of making the plants. The transgenic plants display enhanced resistance to a variety of fungal, bacterial and viral plant pathogens due to expression of a gene that increases the unsaturated fatty acid content of the plant's cells, as compared with an equivalent, but non-transformed plant. The preferred embodiment of the invention is a plant expressing a heterologous DELTA-9 desaturase gene from yeast, which particularly increases cytosolic quantities of 16:1, 16:2 and 18:1 fatty acids.

32. Document ID: US 5874087 A

L9: Entry 32 of 52

File: EPAB

Feb 23, 1999

PUB-NO: US005874087A
DOCUMENT-IDENTIFIER: US 5874087 A
TITLE: Modified plant viruses as vectors

PUBN-DATE: February 23, 1999

INVENTOR-INFORMATION:
NAME

COUNTRY

LOMONOSSOFF, GEORGE PETER

GB

JOHNSON, JOHN EMIL

US

INT-CL (IPC): A61K 39/00; A61K 39/12; C12N 7/01
EUR-CL (EPC): C12N015/82

IN: LOMONOSSOFF, GEORGE PETER, JOHNSON, JOHN EMIL

L9: Entry 32 of 52

File: EPAB

Feb 23, 1999

DOCUMENT-IDENTIFIER: US 5874087 A
TITLE: Modified plant viruses as vectors

FPAR:

The invention relates to assembled particles of a plant virus containing a predetermined foreign peptide as part of the coat protein of the virus, and a method for their production. The foreign peptide is preferably a biologically functional peptide, the biological

application of which
requires or is enhanced by presentation of the peptide in association with a
larger molecule or
particle.

33. Document ID: WO 9856933 A1

L9: Entry 33 of 52

File: EPAB

Dec 17, 1998

PUB-NO: WO009856933A1
DOCUMENT-IDENTIFIER: WO 9856933 A1
TITLE: POLYPEPTIDE PRESENTATION SYSTEM

PUBN-DATE: December 17, 1998

INVENTOR-INFORMATION:
NAME

	COUNTRY
LOMONOSSOFF, GEORGE PETER	
TAYLOR, KATHRYN MAY	GB
	GB

INT-CL (IPC): C12N 15/82; C12N 15/41; C07K 14/095; C12N 7/04;
A61K 39/125

EUR-CL (EPC): C07K014/095; C12N015/82, C12N015/82

IN: LOMONOSSOFF, GEORGE PETER, TAYLOR, KATHRYN
MAY

L9: Entry 33 of 52

File: EPAB

Dec 17, 1998

DOCUMENT-IDENTIFIER: WO 9856933 A1
TITLE: POLYPEPTIDE PRESENTATION SYSTEM

FPAR:

Disclosed are nucleic acid constructs comprising a sequence encoding a
plant viral coat protein
(e.g. the S-peptide of CPMV) containing a foreign or heterologous peptide
insert (e.g. an epitope
for vaccine use) wherein the said coat protein has been modified such as to
reduce its ability to
effect nucleic acid packaging within viral particles. The modification is
preferably at the
C-terminus. Also disclosed are corresponding nucleic acid preparations,
plus methods, processes
and other materials (e.g. plants, virus particles, and compositions) based on
the nucleic acids.

34. Document ID: US 5633434 A

L9: Entry 34 of 52

File: EPAB

May 27, 1997

PUB-NO: US005633434A
DOCUMENT-IDENTIFIER: US 5633434 A
TITLE: Transgenic plants displaying virus and phosphinothricin resistance

PUBN-DATE: May 27, 1997

INVENTOR-INFORMATION:
NAME

	COUNTRY
SCHNEIDER, RUDOLF	
DONN, GUENTER	DE
MUELLNER, HUBERT	DE
	DE

INT-CL (IPC): C12N 5/10; C12N 15/11; C12N 15/33; A01H 5/00

IN: SCHNEIDER, RUDOLF, DONN, GUENTER, MUELLNER,
HUBERT

L9: Entry 34 of 52

File: EPAB

May 27, 1997

DOCUMENT-IDENTIFIER: US 5633434 A
TITLE: Transgenic plants displaying virus and phosphinothricin resistance

35. Document ID: US 5589625 A

L9: Entry 35 of 52

File: EPAB

Dec 31, 1996

PUB-NO: US005589625A
DOCUMENT-IDENTIFIER: US 5589625 A
TITLE: Transgenic plants displaying multiple virus resistance and a process
for their production

PUBN-DATE: December 31, 1996

INVENTOR-INFORMATION:
NAME

	COUNTRY
SAARMA, MART	
KELVE, MERIKKE	FI
TRUVE, ERKKI	EE
TEERI, TEEMU	EE
	FI

INT-CL (IPC): A01H 5/00; C12N 15/82
EUR-CL (EPC): C12N009/12; C12N015/82

IN: SAARMA, MART, KELVE, MERIKKE, TRUVE, ERKKI,
TEERI, TEEMU

L9: Entry 35 of 52

File: EPAB

Dec 31, 1996

DOCUMENT-IDENTIFIER: US 5589625 A
TITLE: Transgenic plants displaying multiple virus resistance and a process for their production

36. Document ID: WO 9640948 A2
L9: Entry 36 of 52
File: EPAB
Dec 19, 1996

PUB-NO: WO009640948A2
DOCUMENT-IDENTIFIER: WO 9640948 A2
TITLE: RESISTANCE TO VIRUSES AND VIROIDS IN TRANSGENIC PLANT AND ANIMAL HOSTS EXPRESSING dsRNA-BINDING PROTEIN

PUBN-DATE: December 19, 1996

INVENTOR-INFORMATION:
NAME

	COUNTRY
ROTH, DON ALLEN	
LANGLAND, JEFFREY OLAF	N/A
	N/A

INT-CL (IPC): C12N 15/82; C12N 15/34; C12N 15/85; A61K 38/16; A61K 48/00; A01N 63/02; A01H 5/00
EUR-CL (EPC): A01K067/027; C07K014/07, C07K014/14, C12N015/82

IN: ROTH, DON ALLEN, LANGLAND, JEFFREY OLAF

L9: Entry 36 of 52
File: EPAB
Dec 19, 1996

DOCUMENT-IDENTIFIER: WO 9640948 A2
TITLE: RESISTANCE TO VIRUSES AND VIROIDS IN TRANSGENIC PLANT AND ANIMAL HOSTS EXPRESSING dsRNA-BINDING PROTEIN

FPAR:

The invention provides a method for imparting resistance in animals to viruses, and in plants to viruses and viroids, that express double-stranded RNA-like structures (dsRNAs). This method enables the binding of pathogenic dsRNAs during the infection process by expression of dsRNA-binding protein in transgenic animal and plant hosts, thus interrupting the infection cycle and inhibiting disease. The presence of a dsRNA-binding protein in a transgenic host renders the transgenic host resistant to the phenotypic symptoms of viral infection and/or decreased pathogen replication. Accordingly, the present invention provides a genetically engineered animal and plant, stably transformed to express a dsRNA-binding protein, such that the transgenic host displays resistance to virus and/or viroid challenge.

37. Document ID: WO 9612028 A1
L9: Entry 37 of 52
File: EPAB
Apr 25, 1996

PUB-NO: WO009612028A1
DOCUMENT-IDENTIFIER: WO 9612028 A1
TITLE: PRODUCTION OF PEPTIDES IN PLANTS AS VIRAL COAT PROTEIN FUSIONS

PUBN-DATE: April 25, 1996

INVENTOR-INFORMATION:
NAME

	COUNTRY
TURPEN, THOMAS H	
REINL, STEPHEN J	N/A
GRILL, LAURENCE K	N/A
	N/A

INT-CL (IPC): C12N 15/82; C12N 15/40; C12N 15/62; C12N 7/01; C12N 5/10
EUR-CL (EPC): C07K014/08; C07K014/445, C12N015/82, C12N015/82, C12N015/82

IN: TURPEN, THOMAS H, REINL, STEPHEN J, GRILL, LAURENCE K

L9: Entry 37 of 52
File: EPAB
Apr 25, 1996

DOCUMENT-IDENTIFIER: WO 9612028 A1
TITLE: PRODUCTION OF PEPTIDES IN PLANTS AS VIRAL COAT PROTEIN FUSIONS

FPAR:

The present invention relates to foreign peptide sequences fused to recombinant plant viral structural proteins and a method of their production. Fusion proteins are economically synthesized in plants at high levels by biologically contained tobamoviruses. The fusion proteins of the invention have many uses. Such uses include use as antigens for inducing the production of antibodies having desired binding properties, e.g., protective antibodies, or for use as vaccine antigens for the induction of protective immunity, including immunity against parasitic infections.

38. Document ID: WO 9602649 A1
L9: Entry 38 of 52
File: EPAB
Feb 1, 1996

PUB-NO: WO009602649A1
DOCUMENT-IDENTIFIER: WO 9602649 A1
TITLE: MODIFIED PLANT VIRUSES AS VECTORS OF HETEROLOGOUS PEPTIDES

PUBN-DATE: February 1, 1996

INVENTOR-INFORMATION:
NAME

COUNTRY

LOMONOSSOFF, GEORGE PETER

GB

INT-CL (IPC): C12N 15/40; C12N 15/41; C12N 15/42; C12N 15/49;
C12N 15/62; C12N 7/01; A61K 39/12
EUR-CL (EPC): C07K014/08; C07K014/09, C07K014/095, C07K014/16
, C12N015/62, C12N015/82,
C12N015/82; C12N015/82

IN: LOMONOSSOFF, GEORGE PETER

L9: Entry 38 of 52

File: EPAB

Feb 1, 1996

DOCUMENT-IDENTIFIER: WO 9602649 A1
TITLE: MODIFIED PLANT VIRUSES AS VECTORS OF
HETEROLOGOUS PEPTIDES

FPAR:
CHG DATE=19990617 STATUS=O>The invention relates to assembled
particles of a plant virus
containing a foreign peptide insert in the coat protein of the virus. The site
of the insert is
free from direct sequence repeats flanking the insert. The invention also
relates to a method of
production of the particles and their use, in particular in vaccines.

39. Document ID: WO 9416087 A1

L9: Entry 39 of 52

File: EPAB

Jul 21, 1994

PUB-NO: WO009416087A1
DOCUMENT-IDENTIFIER: WO 9416087 A1
TITLE: PLANT VIRUS-RESISTANT TRANSGENIC PLANTS AND
METHOD FOR PRODUCING SAME

PUBN-DATE: July 21, 1994

INVENTOR-INFORMATION:
NAME

COUNTRY

LAGAVRE, THIERRY

FR

DURAND-TARDIF, MYLENE

FR

CASSE-DELBART, FRANCINE

FR

ROBAGLIA, CHRISTOPHE

FR

INT-CL (IPC): C12N 15/82; C12N 15/40; C12N 15/57; C12N 15/54;
A01N 63/00; A01H 5/00
EUR-CL (EPC): C07K014/08; C12N009/12, C12N009/50, C12N015/82

IN: LAGAVRE, THIERRY, DURAND-TARDIF, MYLENE,
CASSE-DELBART, FRANCINE, ROBAGLIA,
CHRISTOPHE

L9: Entry 39 of 52

File: EPAB

Jul 21, 1994

DOCUMENT-IDENTIFIER: WO 9416087 A1
TITLE: PLANT VIRUS-RESISTANT TRANSGENIC PLANTS AND
METHOD FOR PRODUCING SAME

FPAR:
A potyvirus-resistant plant having in its genome one or more DNA
fragments expressing transcripts
corresponding to a protein or protein fraction of a donor virus, e.g. a
protein involved in the
formation of a potyvirus replication complex, a protein involved in the
transport of a potyvirus
between cells, or a protein having a similarity with the cleavage sites of
potyvirus viral
proteins. A method for producing plant virus-resistant plants comprises
inserting vectors from a
donor virus complementary cDNA library into cells or tissue fragments
from said plants, and
selecting plants displaying plant virus resistance.

40. Document ID: WO 9319187 A1

L9: Entry 40 of 52

File: EPAB

Sep 30, 1993

PUB-NO: WO009319187A1
DOCUMENT-IDENTIFIER: WO 9319187 A1
TITLE: TRANSGENIC PLANTS DISPLAYING MULTIPLE VIRUS
RESISTANCE AND A PROCESS FOR THEIR PRODUCTION

PUBN-DATE: September 30, 1993

INVENTOR-INFORMATION:
NAME

COUNTRY

SAARMA, MART

FI

KELVE, MERIKKE

@@

TRUVE, ERKKI

@@

TEERI, TEEMU

FI

US-CL-CURRENT: 800/298; 800/FOR.102
INT-CL (IPC): A01H 5/00; A01N 63/00; C12N 15/54; C12N 15/82
EUR-CL (EPC): C12N009/12; C12N015/82
IN: SAARMA, MART, KELVE, MERIKKE, TRUVE, ERKKI,
TEERI, TEEMU

L9: Entry 40 of 52

File: EPAB

Sep 30, 1993

DOCUMENT-IDENTIFIER: WO 9319187 A1
TITLE: TRANSGENIC PLANTS DISPLAYING MULTIPLE VIRUS
RESISTANCE AND A PROCESS FOR THEIR PRODUCTION

FPAR:
CHG DATE=19990617 STATUS=O>This invention relates to transgenic plants displaying multiple virus resistance using parts of the 2,5A oligoadenylate pathway. In particular, said transgenic plants contain a genetically engineered DNA sequence encoding at least one polypeptide having a 2,5A synthetase activity. Moreover, this invention relates to a process for the production of said transgenic plants and to the use of said genetically engineered DNA sequence.

41. Document ID: WO 9218618 A1

L9: Entry 41 of 52

File: EPAB

Oct 29, 1992

PUB-NO: WO009218618A1
DOCUMENT-IDENTIFIER: WO 9218618 A1
TITLE: MODIFIED PLANT VIRUSES AS VECTORS

PUBN-DATE: October 29, 1992

INVENTOR-INFORMATION:
NAME

COUNTRY

LOMONOSSOFF, GEORGE PETER

GB

JOHNSON, JOHN EMIL

US

INT-CL (IPC): A61K 39/12; A61K 39/125; A61K 39/135; A61K 39/21; C12N 7/01; C12N 15/83
EUR-CL (EPC): C12N015/62; C07K014/08, C07K014/09, C07K014/095, C07K014/16

IN: LOMONOSSOFF, GEORGE PETER, JOHNSON, JOHN EMIL

L9: Entry 41 of 52

File: EPAB

Oct 29, 1992

DOCUMENT-IDENTIFIER: WO 9218618 A1
TITLE: MODIFIED PLANT VIRUSES AS VECTORS

FPAR:
The invention relates to assembled particles of a plant virus containing a predetermined foreign peptide as part of the coat protein of the virus, and a method for their production. The foreign peptide is preferably a biologically functional peptide, the biological application of which requires or is enhanced by presentation of the peptide in association with a larger molecule or particle.

42. Document ID: EP 428881 A1

L9: Entry 42 of 52

File: EPAB

May 29, 1991

PUB-NO: EP000428881A1
DOCUMENT-IDENTIFIER: EP 428881 A1
TITLE: RNA with endonuclease and antisense activity, its production and use.

PUBN-DATE: May 29, 1991

INVENTOR-INFORMATION:
NAME

COUNTRY

MUELLNER, HUBERT DR

DE

SCHNEIDER, RUDOLF DR

DE

UHLMANN, EUGEN DR

DE

UIJTEWAAL, BERNARDUS DR

NL

ECKES, PETER DR

DE

INT-CL (IPC): A01H 5/00; A01N 57/16; C12N 5/10; C12N 9/00; C12N 15/11
EUR-CL (EPC): C12N015/11; C12N009/00, C12N015/82

IN: MUELLNER, HUBERT DR, SCHNEIDER, RUDOLF DR, UHLMANN, EUGEN DR, UIJTEWAAL, BERNARDUS DR, ECKES, PETER DR

L9: Entry 42 of 52

File: EPAB

May 29, 1991

DOCUMENT-IDENTIFIER: EP 428881 A1
TITLE: RNA with endonuclease and antisense activity, its production and use.

FPAR:
Host cells can be transformed so that they express ribozyme RNA and antisense RNA, which are linked together in the loop of the ribozyme. The RNA molecules can, for example, be complementary to a particular viral RNA. Plants transformed with genes coding for such RNA display significantly improved defences against viruses.

43. Document ID: EP 421376 A1

L9: Entry 43 of 52

File: EPAB

Apr 10, 1991

PUB-NO: EP000421376A1
DOCUMENT-IDENTIFIER: EP 421376 A1
TITLE: Multifunctional RNA with self-processing activity, its production and use.

PUBN-DATE: April 10, 1991

INVENTOR-INFORMATION:
NAME

MUELLNER, HUBERT DR	COUNTRY	000
UHLMANN, EUGEN DR	DE	C12N015/79
ECKES, PETER DR	DE	WO 200053780 A2
SCHNEIDER, RUDOLF DR	DE	September 14, 2000
UIJTEWAAL, BERNARDUS DR	DE	E
	NL	047
		C12N015/79
		INT-CL (IPC): C12N 15/79; C12N 15/82; C12N 15/86
		IN: DAWSON, W O, LEWANDOWSKI, D J, POGUE, G P, TURPEN, T H
US-CL-CURRENT: 435/183		
INT-CL (IPC): A01H 5/00; A01N 57/16; C12N 5/10; C12N 9/00; C12N 15/11		
EUR-CL (EPC): C12N015/11; C12N009/00, C12N015/82		
		L9: Entry 44 of 52
		File: DWPI
		Sep 28, 2000
IN: MUELLNER, HUBERT DR, UHLMANN, EUGEN DR, ECKES, PETER DR, SCHNEIDER, RUDOLF DR, UIJTEWAAL, BERNARDUS DR		
		DERWENT-ACC-NO: 2000-594328
		DERWENT-WEEK: 200067
		COPYRIGHT 2001 DERWENT INFORMATION LTD
		TITLE: Multiple component RNA vector system for producing foreign RNAs, peptides and proteins in plants, comprises RNA virus-derived RNA replicons and helper viruses
L9: Entry 43 of 52		
	File: EPAB	
		Apr 10, 1991
DOCUMENT-IDENTIFIER: EP 421376 A1		
TITLE: Multifunctional RNA with self-processing activity, its production and use.		
FPAR:		
Host cells can be transformed so that they express ribozyme RNA and antisense RNA which are connected together via a spacer. The RNA molecules can be complementary, for example, to a particular viral RNA. Plants which are transformed with genes coding for such RNA display a significantly improved defence against viruses.		
		45. Document ID: CA 2228730 A1
		L9: Entry 45 of 52
		File: DWPI
		Jun 24, 1999
		DERWENT-ACC-NO: 2000-161486
		DERWENT-WEEK: 200015
		COPYRIGHT 2001 DERWENT INFORMATION LTD
		TITLE: Antimicrobial styelin peptides isolated from <i>Styela clava</i> useful for preserving materials vulnerable to microbial degradation and for protecting plants and animals against pathogenic microbes
44. Document ID: AU 200036183 A, WO 200053780 A2		
L9: Entry 44 of 52		
	File: DWPI	
		Sep 28, 2000
DERWENT-ACC-NO: 2000-594328		
DERWENT-WEEK: 200067		
COPYRIGHT 2001 DERWENT INFORMATION LTD		
TITLE: Multiple component RNA vector system for producing foreign RNAs, peptides and proteins in plants, comprises RNA virus-derived RNA replicons and helper viruses		
INVENTOR: DAWSON, W O; LEWANDOWSKI, D J ; POGUE, G P ; TURPEN, T H		
PRIORITY-DATA: 1999US-0265575 (March 9, 1999)		
PATENT-FAMILY:		
PUB-NO	PUB-DATE	LANGUAGE
		PAGES
		MAIN-IPC
	CA 2228730 A1	
	June 24, 1999	
		E
		035
		C12N015/12
		INT-CL (IPC): A01N 63/02; A23L 3/3544; A61K 7/00; A61K 38/17; A61L 2/16; C07K 14/435; C07K 16/18; C12N 15/12
	IN: LEE, I, LEHRER, R I, ZHAO, C	
AU 200036183 A		
September 28, 2000		
N/A		

L9: Entry 45 of 52

File: DWPI

Jun 24, 1999

DERWENT-ACC-NO: 2000-161486
DERWENT-WEEK: 200015
COPYRIGHT 2001 DERWENT INFORMATION LTD

TITLE: Antimicrobial styelin peptides isolated from *Styela clava* useful for preserving materials vulnerable to microbial degradation and for protecting plants and animals against pathogenic microbes

ABTX:

USE - The compound (I) displays a wide range of antimicrobial activities and are therefore useful for preserving materials susceptible to microbial degradation, for protecting plants against bacterial infection and in the therapeutic and prophylactic protection of animals against bacteria, fungi and viruses. (I) may also be used as standards in antimicrobial assays and as affinity ligands for absorption of counterpart structures in microbes, including viruses.

46. Document ID: HU 200003485 A2, WO 9905319 A2, AU 9885765 A, EP 990047 A2, CN 1270598 A

L9: Entry 46 of 52

File: DWPI

Dec 28, 2000

DERWENT-ACC-NO: 1999-142963
DERWENT-WEEK: 200111
COPYRIGHT 2001 DERWENT INFORMATION LTD

TITLE: New tagged phosphoramidites and phosphonates for labelling nucleic acid - contain group detectable by mass spectrometry, used to detect or quantify polymorphisms or specific RNA, e.g. for genotyping and detecting mutations

INVENTOR: HOWBERT, J; MULLIGAN, J T ; TABONE, J C ; VAN NESS, J

PRIORITY-DATA: 1997US-0898564 (July 22, 1997), 1997US-0898180 (July 22, 1997), 1997US-0898501 (July 22, 1997)

PATENT-FAMILY:

PUB-NO

PUB-DATE

LANGUAGE

PAGES

MAIN-IPC

HU 200003485 A2

December 28, 2000

N/A

000

C07H021/00

WO 9905319 A2

February 4, 1999

E

263

C12Q001/68

AU 9885765 A

February 16, 1999

N/A

000

C12Q001/68

EP 990047 A2

April 5, 2000

E

000

C12Q001/68

CN 1270598 A

October 18, 2000

N/A

000

C07H021/00

INT-CL (IPC): C07H 21/00; C12Q 1/68

IN: HOWBERT, J, MULLIGAN, J T, TABONE, J C, VAN NESS, J

L9: Entry 46 of 52

File: DWPI

Dec 28, 2000

DERWENT-ACC-NO: 1999-142963
DERWENT-WEEK: 200111
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TITLE: New tagged phosphoramidites and phosphonates for labelling nucleic acid - contain group detectable by mass spectrometry, used to detect or quantify polymorphisms or specific RNA, e.g. for genotyping and detecting mutations

ABTX:

USE - Probes and primers derived from (I), i.e. where X is replaced by a nucleic acid sequence, are used as tags for use in any nucleic acid reaction that requires separation of molecules according to size. Typical of many applications are in polymerase chain reactions, differential display, RNA or dideoxy fingerprinting, ligase or nuclease-based assays etc., e.g. in diagnosis (detecting mutations), forensic studies, detecting polymorphisms, genetic mapping (genotyping of animals, plants, bacteria, viruses and fungi), in toxicology, animal breeding, analysis of gene expression and sequencing by hybridisation.

47. Document ID: AU 9880300 A, WO 9856933 A1

L9: Entry 47 of 52

File: DWPI

Dec 30, 1998

DERWENT-ACC-NO: 1999-060334
DERWENT-WEEK: 199920
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TITLE: New nucleic acid constructs encoding a modified plant viral protein - useful for provoking an immune response in a mammal

INVENTOR: LOMONOSOFF, G P; TAYLOR, K M

PRIORITY-DATA: 1997GB-0012282 (June 12, 1997)

PATENT-FAMILY:

PUB-NO

PUB-DATE

LANGUAGE
PAGES
MAIN-IPC
AU 9880300 A
December 30, 1998
N/A
000
C12N015/82
WO 9856933 A1
December 17, 1998
E
047
C12N015/82
INT-CL (IPC): A61K 39/125; C07K 14/095; C12N 7/04; C12N 15/41;
C12N 15/82

IN: LOMONOSSOFF, G P, TAYLOR, K M

L9: Entry 47 of 52

File: DWPI

Dec 30, 1998

DERWENT-ACC-NO: 1999-060334
DERWENT-WEEK: 199920
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TITLE: New nucleic acid constructs encoding a modified plant viral
protein - useful for provoking
an immune response in a mammal

ABTX:

Also claimed are: (1) a nucleic acid composition (I) containing sequences
which encode other
components required for assembly and/or replication of viral particles. The
peptide insert is
displayed on their surface; (2) a host cell containing (1), preferably a cow
pea plant cell; (3)
a part of a plant containing modified viral particles; and (4) A population of
modified plant
virus particles.

48. Document ID: EP 979243 A2, WO 9841535 A2, AU 9865098
A

L9: Entry 48 of 52

File: DWPI

Feb 16, 2000

DERWENT-ACC-NO: 1998-521161
DERWENT-WEEK: 200014
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TITLE: New modified peptide(s) - obtained by substitution with an amino
acid which is modifiable
by a reaction and replacing other amino acids which are not to be modified

INVENTOR: AJOULA, H S; CLARKE, D J

PRIORITY-DATA: 1997GB-0005519 (March 18, 1997)

PATENT-FAMILY:
PUB-NO

PUB-DATE

LANGUAGE
PAGES
MAIN-IPC

EP 979243 A2
February 16, 2000
E
000
C07K007/08
WO 9841535 A2
September 24, 1998
E
033
C07K000/00
AU 9865098 A
October 12, 1998
N/A
000
C07K001/00

INT-CL (IPC): A61K 38/10; A61K 38/17; C07K 0/00; C07K 1/00; C07K
7/08; C07K 14/435; G01N 33/68

IN: AJOULA, H S, CLARKE, D J

L9: Entry 48 of 52

File: DWPI

Feb 16, 2000

DERWENT-ACC-NO: 1998-521161
DERWENT-WEEK: 200014
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TITLE: New modified peptide(s) - obtained by substitution with an amino
acid which is modifiable
by a reaction and replacing other amino acids which are not to be modified

ABTX:

USE - The methods can be used for the modification of biologically active
peptides such as
hormones, drugs, toxins and peptides which act on lipid bilayer
membranes. The modified peptides
can be used e.g. in the body of an animal or plant or parts in order to affect
the structure or
integrity or permeability of a foreign body such as a microorganism,
parasite or virus present in
the body of the animal or plant or within the cells of the body of the animal
or plant. They can
also be used in assays, purifications or sensors.

49. Document ID: WO 9602649 A1, AU 9528934 A, CZ 9600723
A3, EP 719336 A1, SK 9600342 A3, JP
09504961 W, BR 9506047 A, HU 75088 T, NZ 289170 A, CN 1134174
A, AU 697867 B, US 5958422 A

L9: Entry 49 of 52

File: DWPI

Feb 1, 1996

DERWENT-ACC-NO: 1996-105911
DERWENT-WEEK: 199948
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TITLE: Plant virus assembled particles contg. foreign peptide insert - useful
in vaccines, esp.
for protecting animals, including humans, from virus infections

INVENTOR: LOMONOSSOFF, G P

PRIORITY-DATA: 1994GB-0014118 (July 13, 1994)

PATENT-FAMILY:
PUB-NO

Feb 1, 1996

PATENT-FAMILY: PUB-NO	PUB-DATE	LANGUAGE PAGES MAIN-IPC
WO 9602649 A1	February 1, 1996	E 043 C12N015/40
AU 9528934 A	February 16, 1996	N/A 000 C12N015/40
CZ 9600723 A3	June 12, 1996	N/A 000 C12N015/40
EP 719336 A1	July 3, 1996	E 000 C12N015/40
SK 9600342 A3	July 3, 1996	N/A 000 C12N015/40
JP 09504961 W	May 20, 1997	N/A 038 C12N015/09
BR 9506047 A	August 5, 1997	N/A 000 C12N015/40
HU 75088 T	April 28, 1997	N/A 000 C12N015/40
NZ 289170 A	November 24, 1997	N/A 000 C12N007/01
CN 1134174 A	October 23, 1996	N/A 000 C12N015/40
AU 697867 B	October 22, 1998	N/A 000 C12N015/40
US 5958422 A	September 28, 1999	N/A 000 A61K039/12

INT-CL (IPC): A01H 1/00; A61K 39/00; A61K 39/12; C07H 21/04; C12N 5/00; C12N 5/10; C12N 7/00; C12N 7/01; C12N 15/09; C12N 15/40; C12N 15/41; C12N 15/42; C12N 15/49; C12N 15/62; C12N 15/64

IN: LOMONOSOFF, G P

L9: Entry 49 of 52

File: DWPI

DERWENT-ACC-NO: 1996-105911
DERWENT-WEEK: 199948
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TITLE: Plant virus assembled particles contg. foreign peptide insert -
useful in vaccines, esp.
for protecting animals, including humans, from virus infections

ABTX:

Assembled particles of a plant virus, contg. a foreign peptide insert (A) in the coat protein region, are new. The site of the insert is free of direct sequence repeats flanking the insert, and the virus is pref. Cowpea Mosaic Virus (CPMV). Also claimed are: (1) a fragment of CPMV coat protein cDNA, contg. DNA encoding (A) or a site corresp. to an exposed surface of a coat protein, free from direct repeats flanking the insert; (2) a vector contg. the fragment of (1) which is esp. a full-length cDNA of CPMV M RNA contg. the foreign DNA insert; and (3) an RNA transcript of (1) or (2), which is pref. capped.

ABEQ:

Assembled particles of a plant virus, contg. a foreign peptide insert (A) in the coat protein region, are new. The site of the insert is free of direct sequence repeats flanking the insert, and the virus is pref. Cowpea Mosaic Virus (CPMV). Also claimed are: (1) a fragment of CPMV coat protein cDNA, contg. DNA encoding (A) or a site corresp. to an exposed surface of a coat protein, free from direct repeats flanking the insert; (2) a vector contg. the fragment of (1) which is esp. a full-length cDNA of CPMV M RNA contg. the foreign DNA insert; and (3) an RNA transcript of (1) or (2), which is pref. capped.

TTX:

PLANT VIRUS ASSEMBLE PARTICLE CONTAIN FOREIGN
PEPTIDE INSERT USEFUL VACCINE PROTECT ANIMAL HUMAN
VIRUS INFECT

50. Document ID: ES 2155062 T3, WO 9319187 A1, AU 9226613 A, FI 9404315 A, NO 9403439 A, EP 632835 A1, JP 07504820 W, HU 70265 T, BR 9207103 A, AU 669130 B, US 5589625 A, RU 2125606 C1, HU 218356 B, EP 632835 B1, DE 69231711 E

L9: Entry 50 of 52

File: DWPI

May 1, 2001

DERWENT-ACC-NO: 1993-320753
DERWENT-WEEK: 200136
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TITLE: Transgenic plants with multiple virus resistance - genetically
engineered using DNA
encoding 2,5A synthetase

INVENTOR: KELVE, M; SAARMA, M; TEERI, T; TRUVE, E

PRIORITY-DATA: 1992EP-0104676 (March 18, 1992)

PATENT-FAMILY:
PUB-NO

PUB-DATE	LANGUAGE	PAGES	MAIN-IPC
ES 2155062 T3 May 1, 2001	N/A	000	C12N015/82
WO 9319187 A1 September 30, 1993	E	024	C12N015/82
AU 9226613 A October 21, 1993	N/A	000	C12N015/82
FI 9404315 A September 16, 1994	N/A	000	A01H000/00
NO 9403439 A November 11, 1994	N/A	000	C12N015/82
EP 632835 A1 January 11, 1995	E	000	C12N015/82
JP 07504820 W June 1, 1995	N/A	000	A01H005/00
HU 70265 T September 28, 1995	N/A	000	C12N015/82
BR 9207103 A December 19, 1995	N/A	000	C12N015/82
AU 669130 B May 30, 1996	N/A	000	C12N015/54
US 5589625 A December 31, 1996	N/A	017	A01H005/00
RU 2125606 C1 January 27, 1999	N/A	000	C12N015/82
HU 218356 B August 28, 2000	N/A	000	C12N015/82
EP 632835 B1 February 28, 2001	E	000	C12N015/82
DE 69231711 E April 5, 2001	N/A	000	

C12N015/82

INT-CL (IPC): A01H 0/00; A01H 1/00; A01H 5/00; A01N 63/00; C12N 5/10; C12N 15/09; C12N 15/12; C12N 15/54; C12N 15/82

IN: KELVE, M, SAARMA, M, TEERI, T, TRUVE, E

L9: Entry 50 of 52

File: DWPI

May 1, 2001

DERWENT-ACC-NO: 1993-320753
DERWENT-WEEK: 200136
COPYRIGHT 2001 DERWENT INFORMATION LTD

TITLE: Transgenic plants with multiple virus resistance - genetically engineered using DNA encoding 2,5A synthetase

AB TX:

A transgenic plant (I) displaying multiple virus resistance contains a genetically engineered DNA sequence encoding at least one polypeptide with 2,5A synthetase activity. The polypeptide is capable of activating an endonuclease causing degradation of viral RNA.

ABEQ:

A transgenic plant (I) displaying multiple virus resistance contains a genetically engineered DNA sequence encoding at least one polypeptide with 2,5A synthetase activity. The polypeptide is capable of activating an endonuclease causing degradation of viral RNA.

ABEQ:

A transgenic plant that utilizes the 2,5A oligoadenylate pathway and displays resistance to multiple plant viral taxonomic groups comprising a genetically engineered DNA sequence encoding at least one polypeptide having a 2,5A synthetase activity, wherein said polypeptide activates an endonuclease causing degradation of viral RNA.